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[YU/US]; 1006 Asbury Way, Mountain View, CA 94043 (US).

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- (74) Agents: FRANCIS, Carol, L. et al.: Bozicevic, Field & Francis LLP, 200 Middlefield Road, Suite 200. Menlo Park, CA 94025 (US).

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- (71) Applicants (for all designated States except US): CHI-RON CORPORATION [US/US]: 4560 Horton Street. Emeryville, CA 94608-2916 (US). HYSEQ INC. [US/US]; 675 Almanor Avenue, Sunnyvale, CA 94086 (US).
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(72) Inventors; and

Published:

(75) Inventors/Applicants (for US only): ESCOBEDO, Jaime [CL/US]; 1470 Lavorna Road, Alamo, CA 94507 (US), GARCIA, Pablo, Dominguez [CL/US]; 882 Chenery Street, San Francisco, CA 94131 (US). SUD-DUTH-KLINGER, Julie [US/US]; 280 Lexington Road, Kensington, CA 94707 (US). REINHARD, Christoph [DE/US]; 1633 Clinton Avenue, Alameda, CA 94504 (US). RANDAZZO, Filippo [US/US]; 104 Capricorn Avenue, Oakland, CA 94611 (US), LAMSON, George [US/US]; 232 Sandringham Drive, Moraga, CA 94556 (US). SCOTT, Elizabeth, M. [US/US]; 1267 Nash Street, Sonoma, CA 95476 (US). ZHANG, Guozhong [CN/US]; 41236 Norman Court, Fremont, CA 94539 (US), KAS-SAM, Altaf [US/US]; 3810 Midvale Avenue, Oakland, CA 94602 (US). POT, David [CA/US]; 5550 Columbia

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(54) Title: HUMAN GENES AND GENE EXPRESSION PRODUCTS

(57) Abstract: This invention relates to novel human polynucleotides and variants thereof, their encoded polypeptides and variants thereof, to genes corresponding to these polynucleotides and to proteins expressed by the genes. The invention also relates to diagnostic and therapeutic agents employing such novel human polynucleotides, their corresponding genes or gene products, e.g., these genes and proteins, including probes, antisense constructs, and antibodies.

HUMAN GENES AND GENE EXPRESSION PRODUCTS

Cross-Reference to Related Application

[0001] The application claims the benefit of earlier-filed U.S. provisional application serial no. 60/226,326 filed August 16, 2000, which application is incorporated herein by reference in its entirety.

Field of the Invention

[0002] The present invention relates to polynucleotides of human origin, particularly in human colon, breast, prostate, and/or lung tissue, and the encoded gene products.

Background of the Invention

[0003] Identification of novel polynucleotides, particularly those that encode an expressed gene product, is important in the advancement of drug discovery, diagnostic technologies, and the understanding of the progression and nature of complex diseases such as cancer. Identification of genes expressed in different cell types isolated from sources that differ in disease state or stage, developmental stage, exposure to various environmental factors, the tissue of origin, the species from which the tissue was isolated, and the like is key to identifying the genetic factors that are responsible for the phenotypes associated with these various differences.

[0004] This invention provides novel human polynucleotides, the polypeptides encoded by these polynucleotides, and the genes and proteins corresponding to these novel polynucleotides. Summary of the Invention

[0005] This invention relates to novel human polynucleotides and variants thereof, their encoded polypeptides and variants thereof, to genes corresponding to these polynucleotides and to proteins expressed by the genes. The invention also relates to diagnostics and therapeutics comprising such novel human polynucleotides, their corresponding genes or gene products, including probes, antisense nucleotides, and antibodies. The polynucleotides of the invention correspond to a polynucleotide comprising the sequence information of at least one of SEQ ID NOS:1-6010.

[0006] Various aspects and embodiments of the invention will be readily apparent to the ordinarily skilled artisan upon reading the description provided herein.

Detailed Description of the Invention

[0007] Before the present invention is described, it is to be understood that this invention is not limited to particular embodiments described, as such may, of course, vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodiments only, and is not intended to be limiting.

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Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. Although any methods and materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, the preferred methods and materials are now described.

All publications and patent applications cited in this specification are herein incorporated by reference as if each individual publication or patent application were specifically and individually indicated to be incorporated by reference. The citation of any publication is for its disclosure prior to the filing date and should not be construed as an admission that the present invention is not entitled to antedate such publication by virtue of prior invention.

It must be noted that as used herein and in the appended claims, the singular forms "a," "and," and "the" include plural referents unless the context clearly dictates otherwise. Thus, for example, reference to "a polynucleotide" includes a plurality of such polynucleotides and reference to "the colon cancer cell" includes reference to one or more cells and equivalents thereof known to those skilled in the art, and so forth.

The publications and applications discussed herein are provided solely for their disclosure prior to the filing date of the present application. Nothing herein is to be construed as an admission that the present invention is not entitled to antedate such publication by virtue of prior invention. Further, the dates of publication provided may be different from the actual publication dates which may need to be independently confirmed.

Definitions

The terms "polynucleotide" and "nucleic acid," used interchangeably herein, refer to a polymeric forms of nucleotides of any length, either ribonucleotides or deoxynucleotides. Thus, these terms include, but are not limited to, single-, double-, or multi-stranded DNA or RNA, genomic DNA, cDNA, DNA-RNA hybrids, branched nucleic acid (see, e.g., U.S. Pat. Nos. 5,124,246; 5,710,264; and 25 5,849,481), or a polymer comprising purine and pyrimidine bases or other natural, chemically or biochemically modified, non-natural, or derivatized nucleotide bases. These terms further include, but are not limited to, mRNA or cDNA that comprise intronic sequences (see, e.g., Niwa et al. (1999) Cell 99(7):691-702). The backbone of the polynucleotide can comprise sugars and phosphate groups (as may typically be found in RNA or DNA), or modified or substituted sugar or phosphate groups. 30 Alternatively, the backbone of the polynucleotide can comprise a polymer of synthetic subunits such as phosphoramidites and thus can be an oligodeoxynucleoside phosphoramidate or a mixed phosphoramidate-phosphodiester oligomer. Peyrottes et al. (1996) Nucl. Acids Res. 24:1841-1848; Chaturvedi et al. (1996) Nucl. Acids Res. 24:2318-2323. A polynuclotide may comprise modified nucleotides, such as methylated nucleotides and nucleotide analogs, uracyl, other sugars, and linking

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groups such as fluororibose and thioate, and nucleotide branches. The sequence of nucleotides may be interrupted by non-nucleotide components. A polynucleotide may be further modified after polymerization, such as by conjugation with a labeling component. Other types of modifications included in this definition are caps, substitution of one or more of the naturally occurring nucleotides with an analog, and introduction of means for attaching the polynucleotide to proteins, metal ions, labeling components, other polynucleotides, or a solid support.

The terms "polypeptide" and "protein," used interchangebly herein, refer to a polymeric form of amino acids of any length, which can include coded and non-coded amino acids, chemically or biochemically modified or derivatized amino acids, and polypeptides having modified peptide backbones. The term includes fusion proteins, including, but not limited to, fusion proteins with a heterologous amino acid sequence, fusions with heterologous and homologous leader sequences, with or without N-terminal methionine residues; immunologically tagged proteins; and the like.

"Diagnosis" as used herein generally includes determination of a subject's susceptibility to a disease or disorder, determination as to whether a subject is presently affected by a disease or disorder, prognosis of a subject affected by a disease or disorder (e.g., identification of pre-metastatic or metastatic cancerous states, stages of cancer, or responsiveness of cancer to therapy), and therametrics (e.g., monitoring a subject's condition to provide information as to the effect or efficacy of therapy).

"Sample" or "biological sample" as used herein encompasses a variety of sample types, and are generally meant to refer to samples of biological fluids or tissues, particularly samples obtained from tissues, especially from cells of the type associated with a disease or condition for which a diagnostic application is designed (e.g., ductal adenocarcinoma), and the like. "Sample" or "biological sample" are meant to encompass blood and other liquid samples of biological origin, solid tissue samples, such as a biopsy specimen or tissue cultures or cells derived therefrom and the progeny thereof. These terms encompass samples that have been manipulated in any way after their procurement as well as derivatives and fractions of samples, where the samples may be manipulated by, for example, treatment with reagents, solubilization, or enrichment for certain components. The terms also encompass clinical samples, and also includes cells in cell culture, cell supernatants, cell lysates, serum, plasma, biological fluids, and tissue samples. Where the sample is solid tissue, the cells of the tissue can be dissociated or tissue sections can be analyzed.

The terms "treatment," "treating," "treat" and the like are used herein to generally refer to obtaining a desired pharmacologic and/or physiologic effect. The effect may be prophylactic in terms of completely or partially preventing a disease or symptom thereof and/or may be therapeutic in terms of a partial or complete stabilization or cure for a disease and/or adverse effect attributable to the disease. "Treatment" as used herein covers any treatment of a disease in a mammal, particularly a

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human, and includes: (a) preventing the disease or symptom from occurring in a subject which may be predisposed to the disease or symptom but has not yet been diagnosed as having it; (b) inhibiting the disease symptom, i.e., arresting its development; or relieving the disease symptom, i.e., causing regression of the disease or symptom.

The terms "individual," "subject," "host," and "patient," used interchangeably herein and refer to any mammalian subject for whom diagnosis, treatment, or therapy is desired, particularly humans. Other subjects may include cattle, dogs, cats, guinea pigs, rabbits, rats, mice, horses, and so on.

As used herein the term "isolated" refers to a polymucleotide, a polypeptide, an antibody, or a host cell that is in an environment different from that in which the polymucleotide, the polypeptide, the antibody, or the host cell naturally occurs. A polymucleotide, a polypeptide, an antibody, or a host cell which is isolated is generally substantially purified. As used herein, the term "substantially purified" refers to a compound (e.g., either a polymucleotide or a polypeptide or an antibody) that is removed from its natural environment and is at least 60% free, preferably 75% free, and most preferably 90% free from other components with which it is naturally associated. Thus, for example, a composition containing A is "substantially free of" B when at least 85% by weight of the total A+B in the composition is A. Preferably, A comprises at least about 90% by weight of the total of A+B in the composition, more preferably at least about 95% or even 99% by weight.

A "host cell," as used herein, refers to a microorganism or a eukaryotic cell or cell line cultured as a unicellular entity which can be, or has been, used as a recipient for a recombinant vector or other transfer polynucleotides, and include the progeny of the original cell which has been transfected. It is understood that the progeny of a single cell may not necessarily be completely identical in morphology or in genomic or total DNA complement as the original parent, due to natural, accidental, or deliberate mutation

The terms "cancer," "neoplasm," "tumor," and "carcinoma," are used interchangeably herein to refer to cells which exhibit relatively autonomous growth, so that they exhibit an aberrant growth phenotype characterized by a significant loss of control of cell proliferation. In general, cells of interest for detection or treatment in the present application include precancerous (e.g., benign), malignant, metastatic, and non-metastatic cells. Detection of cancerous cell is of particular interest.

The use of "e", as in 10e-3, indicates that the number to the left of "e" is raised to the power of the number to the right of "e" (thus, 10e-3 is 10⁻³).

The term "heterologous" as used herein in the context of, for example, heterologous nucleic acid or amino acid sequences, heterologous polypeptides, or heterologous nucleic acid, is meant to refer to material that originates from a source different from that with which it is joined or associated. For example, two DNA sequences are heterologous to one another if the sequences are from different genes

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or from different species. A recombinant host cell containing a sequence that is heterologous to the host cell can be, for example, a bacterial cell containing a sequence encoding a human polypeptide.

The invention relates to polynucleotides comprising the disclosed nucleotide sequences, to full length cDNA, mRNA, genomic sequences, and genes corresponding to these sequences and degenerate variants thereof, and to polypeptides encoded by the polynucleotides of the invention and polypeptide variants. The following detailed description describes the polynucleotide compositions encompassed by the invention, methods for obtaining cDNA or genomic DNA encoding a full-length gene product, expression of these polynucleotides and genes, identification of structural motifs of the polynucleotides and genes, identification of the function of a gene product encoded by a gene corresponding to a polynucleotide of the invention, use of the provided polynucleotides as probes and in mapping and in tissue profiling, use of the corresponding polypeptides and other gene products to raise antibodies, and use of the polynucleotides and their encoded gene products for therapeutic and diagnostic purposes. Polynucleotide Compositions

The scope of the invention with respect to polynucleotide compositions includes, but is not necessarily limited to, polynucleotides having a sequence set forth in any one of SEQ ID NOS:1-6010; polynucleotides obtained from the biological materials described herein or other biological sources (particularly human sources) by hybridization under stringent conditions (particularly conditions of high stringency); genes corresponding to the provided polynucleotides; variants of the provided polynucleotides and their corresponding genes, particularly those variants that retain a biological activity of the encoded gene product (e.g., a biological activity ascribed to a gene product corresponding to the provided polynucleotides as a result of the assignment of the gene product to a protein family(ies) and/or identification of a functional domain present in the gene product). Other nucleic acid compositions contemplated by and within the scope of the present invention will be readily apparent to one of ordinary skill in the art when provided with the disclosure here. "Polynucleotide" and "nucleic acid" as used herein with reference to nucleic acids of the composition is not intended to be limiting as to the length or structure of the nucleic acid unless specifically indicated.

The invention features polynucleotides that are expressed in human tissue, especially human colon, prostate, breast, lung and/or endothelial tissue. Novel nucleic acid compositions of the invention of particular interest comprise a sequence set forth in any one of SEQ ID NOS:1-6010 or an identifying sequence thereof. An "identifying sequence" is a contiguous sequence of residues at least about 10 nt to about 20 nt in length, usually at least about 50 nt to about 100 nt in length, that uniquely identifies a polynucleotide sequence, e.g., exhibits less than 90%, usually less than about 80% to about 85% sequence identity to any contiguous nucleotide sequence of more than about 20 nt. Thus, the

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subject novel nucleic acid compositions include full length cDNAs or mRNAs that encompass an identifying sequence of contiguous nucleotides from any one of SEQ ID NOS: 1-6010.

The polynucleotides of the invention also include polynucleotides having sequence similarity or sequence identity. Nucleic acids having sequence similarity are detected by hybridization under low stringency conditions, for example, at 50°C and 10XSSC (0.9 M saline/0.09 M sodium cirrate) and remain bound when subjected to washing at 55°C in 1XSSC. Sequence identity can be determined by hybridization under stringent conditions, for example, at 50°C or higher and 0.1XSSC (9 mM saline/0.9 mM sodium citrate). Hybridization methods and conditions are well known in the art, see, e.g., USPN 5,707,829. Nucleic acids that are substantially identical to the provided polynucleotide sequences, e.g. allelic variants, genetically altered versions of the gene, etc., bind to the provided polynucleotide sequences (SEQ ID NOS:1-6010) under stringent hybridization conditions. By using probes, particularly labeled probes of DNA sequences, one can isolate homologous or related genes. The source of homologous genes can be any species, e.g. primate species, particularly human; rodents, such as rats and mice; canines, felines, bovines, ovines, equines, yeast, nematodes, etc.

Preferably, hybridization is performed using at least 15 contiguous nucleotides (nt) of at least one of SEQ ID NOS:1-6010. That is, when at least 15 contiguous nt of one of the disclosed SEQ ID NOS. is used as a probe, the probe will preferentially hybridize with a nucleic acid comprising the complementary sequence, allowing the identification and retrieval of the nucleic acid identification the selected probe. Probes from more than one SEQ ID NO. can hybridize with the same nucleic acid if the cDNA from which they were derived corresponds to one mRNA. Probes of more than 15 nt can be used, e.g., probes of from about 18 nt to about 100 nt, but 15 nt represents sufficient sequence for unique identification.

The polynucleotides of the invention also include naturally occurring variants of the nucleotide sequences (e.g., degenerate variants, allelic variants, etc.). Variants of the polynucleotides of the invention are identified by hybridization of putative variants with nucleotide sequences disclosed herein, preferably by hybridization under stringent conditions. For example, by using appropriate wash conditions, variants of the polynucleotides of the invention can be identified where the allelic variant exhibits at most about 25-30% base pair (bp) mismatches relative to the selected polynucleotide probe. In general, allelic variants contain 15-23% bp mismatches, and can contain as little as even 5-15%, or 1-2% bp mismatches, as well as a single bp mismatch.

The invention also encompasses homologs corresponding to the polynucleotides of SEQ ID NOS:1-6010, where the source of homologous genes can be any mammalian species, e.g., primate species, particularly human, rodents, such as rats; canines, felines, bovines, ovines, equines, yeast, nematodes, etc. Between mammalian species, e.g., human and mouse, homologs generally have

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substantial sequence similarity, e.g., at least 75% sequence identity, usually at least 90%, more usually at least 95% between nucleotide sequences. Sequence similarity is calculated based on a reference sequence, which may be a subset of a larger sequence, such as a conserved moit, coding region, flanking region, etc. A reference sequence will usually be at least about 18 contiguous nt long, more usually at least about 30 nt long, and may extend to the complete sequence that is being compared. Algorithms for sequence analysis are known in the art, such as gapped BLAST, described in Altschul, et al. Nuclete Acids Res. (1997) 25:3389-3402.

In general, variants of the invention have a sequence identity greater than at least about 65%, preferably at least about 75%, more preferably at least about 85%, and can be greater than at least about 90% or more as determined by the Smith-Waterman homology search algorithm as implemented in MPSRCH program (Oxford Molecular). For the purposes of this invention, a preferred method of calculating percent identity is the Smith-Waterman algorithm, using the following. Global DNA sequence identity must be greater than 65% as determined by the Smith-Waterman homology search algorithm as implemented in MPSRCH program (Oxford Molecular) using an affine gap search with the following search parameters: gap open penalty, 12; and gap extension penalty, 1.

The subject nucleic acids can be cDNAs or genomic DNAs, as well as fragments thereof, particularly fragments that encode a biologically active gene product and/or are useful in the methods disclosed herein (e.g., in diagnosis, as a unique identifier of a differentially expressed gene of interest, etc.). The term "cDNA" as used herein is intended to include all nucleic acids that share the arrangement of sequence elements found in native mature mRNA species, where sequence elements are exons and 3' and 5' non-coding regions. Normally mRNA species have contiguous exons, with the intervening introns, when present, being removed by nuclear RNA splicing, to create a continuous open reading frame encoding a polyveptide of the invention.

A genomic sequence of interest comprises the nucleic acid present between the initiation codon and the stop codon, as defined in the listed sequences, including all of the introns that are normally present in a native chromosome. It can further include the 3' and 5' untranslated regions found in the mature mRNA. It can further include specific transcriptional and translational regulatory sequences, such as promoters, enhancers, etc., including about 1 kb, but possibly more, of flanking genomic DNA at either the 5' and 3' end of the transcribed region. The genomic DNA can be isolated as a fragment of 100 kbp or smaller; and substantially free of flanking chromosomal sequence. The genomic DNA flanking the coding region, either 3' and 5', or internal regulatory sequences as sometimes found in introns, contains sequences required for proper tissue, stage-specific, or disease-state specific expression.

The nucleic acid compositions of the subject invention can encode all or a part of the subject polypeptides. Double or single stranded fragments can be obtained from the DNA sequence by chemically synthesizing oligonucleotides in accordance with conventional methods, by restriction enzyme digestion, by PCR amplification, etc. Isolated polynucleotides and polynucleotide fragments of the invention comprise at least about 10, about 15, about 20, about 35, about 50, about 100, about 150 to about 200, about 250 to about 300, or about 350 contiguous in selected from the polynucleotide sequences as shown in SEQ ID NOS:1-6010. For the most part, fragments will be of at least 15 nt, usually at least 18 nt or 25 nt, and up to at least about 50 contiguous nt in length or more. In a preferred embodiment, the polynucleotide molecules comprise a contiguous sequence of at least 12 nt selected from the group consisting of the polynucleotides shown in SEO ID NOS:1-6010.

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Probes specific to the polynucleotides of the invention can be generated using the polynucleotide sequences disclosed in SEQ ID NOS:1-6010. The probes are preferably at least about 12, 15, 16, 18, 20, 22, 24, or 25 nt fragment of a corresponding contiguous sequence of SEQ ID NOS:1-6010, and can be less than 2, 1, 0.5, 0.1, or 0.05 kb in length. The probes can be synthesized chemically or can be generated from longer polynucleotides using restriction enzymes. The probes can be labeled, for example, with a radioactive, biotinylated, or fluorescent tag. Preferably, probes are designed based upon an identifying sequence of a polynucleotide of one of SEQ ID NOS:1-6010. More preferably, probes are designed based on a contiguous sequence of one of the subject polynucleotides that remain unmasked following application of a masking program for masking low complexity (e.g., XBLAST) to the sequence, i.e., one would select an unmasked region, as indicated by the polynucleotides outside the poly-n stretches of the masked sequence produced by the masking program.

The polynucleotides of the subject invention are isolated and obtained in substantial purity, generally as other than an intact chromosome. Usually, the polynucleotides, either as DNA or RNA, will be obtained substantially free of other naturally-occurring nucleic acid sequences, generally being at least about 50%, usually at least about 90% pure and are typically "recombinant," e.g., flanked by one or more nucleotides with which it is not normally associated on a naturally occurring chromosome.

The polynucleotides of the invention can be provided as a linear molecule or within a circular molecule, and can be provided within autonomously replicating molecules (vectors) or within molecules without replication sequences. Expression of the polynucleotides can be regulated by their own or by other regulatory sequences known in the art. The polynucleotides of the invention can be introduced into suitable host cells using a variety of techniques available in the art, such as transferrin polycation-mediated DNA transfer, transfertion with naked or encapsulated nucleic acids, liposome-

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mediated DNA transfer, intracellular transportation of DNA-coated latex beads, protoplast fusion, viral infection, electroporation, gene gun, calcium phosphate-mediated transfection, and the like.

The subject nucleic acid compositions can be used to, for example, produce polypeptides, as probes for the detection of mRNA of the invention in biological samples (e.g., extracts of human cells) to generate additional copies of the polynucleotides, to generate ribozymes or antisense oligonucleotides, and as single stranded DNA probes or as triple-strand forming oligonucleotides. The probes described herein can be used to, for example, determine the presence or absence of the polynucleotide sequences as shown in SEQ ID NOS:1-6010 or variants thereof in a sample. These and other uses are described in more detail below.

10 Use of Polynucleotides to Obtain Full-Length cDNA, Gene, and Promoter Region

In one embodiment, the polynucleotides are useful as starting materials to construct larger molecules. In one example, the polynucleotides of the invention are used to construct polynucleotides that encode a larger polypeptide (e.g., up to the full-length native polypeptide as well as fusion proteins comprising all or a portion of the native polypeptide) or may be used to produce haptens of the polypeptide (e.g., polypeptides useful to generate antibodies).

In one particular example, the polynucleotides of the invention are used to make or isolate cDNA molecules encoding all or portion of a naturally-occuring polypeptide. Full-length cDNA molecules comprising the disclosed polynucleotides are obtained as follows. A polynucleotide having a sequence of one of SEQ ID NOS:1-6010, or a portion thereof comprising at least 12, 15, 18, or 20 nt. is used as a hybridization probe to detect hybridizing members of a cDNA library using probe design methods, cloning methods, and clone selection techniques such as those described in USPN 5,654,173. Libraries of cDNA are made from selected tissues, such as normal or tumor tissue, or from tissues of a mammal treated with, for example, a pharmaceutical agent. Preferably, the tissue is the same as the tissue from which the polynucleotides of the invention were isolated, as both the polynucleotides described herein and the cDNA represent expressed genes. Most preferably, the cDNA library is made from the biological material described herein in the Examples. The choice of cell type for library construction can be made after the identity of the protein encoded by the gene corresponding to the polymicleotide of the invention is known. This will indicate which tissue and cell types are likely to express the related gene, and thus represent a suitable source for the mRNA for generating the cDNA. Where the provided polynucleotides are isolated from cDNA libraries, the libraries are prepared from mRNA of human colon cells, more preferably, human colon cancer cells, even more preferably, from a highly metastatic colon cell, Km12L4-A.

Techniques for producing and probing nucleic acid sequence libraries are described, for example, in Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd Ed., (1989) Cold Spring

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Harbor Press, Cold Spring Harbor, NY. The cDNA can be prepared by using primers based on polynucleotides comprising a sequence of SEQ ID NOS:1-6010. In one embodiment, the cDNA library can be made from only poly-adenylated mRNA. Thus, poly-T primers can be used to prepare cDNA from the mRNA.

Members of the library that are larger than the provided polynucleotides, and preferably that encompass the complete coding sequence of the native message, are obtained. In order to confirm that the entire cDNA has been obtained, RNA protection experiments are performed as follows.

Hybridization of a full-length cDNA to an mRNA will protect the RNA from RNase degradation. If the cDNA is not full length, then the portions of the mRNA that are not hybridized will be subject to RNase degradation. This is assayed, as is known in the art, by changes in electrophoretic mobility on polyacrylamide gels, or by detection of released monoribonucleotides. Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd Ed., (1989) Cold Spring Harbor Press, Cold Spring Harbor, NY. In order to obtain additional sequences 5't ot the end of a partial cDNA, 5' RACE (PCR Protecols: A Guide to Methods and Applications, (1990) Academic Press, Inc.) can be performed.

Genomic DNA is isolated using the provided polynucleotides in a manner similar to the isolation of full-length cDNAs. Briefty, the provided polynucleotides, or portions thereof, are used as probes to libraries of genomic DNA. Preferably, the library is obtained from the cell type that was used to generate the polynucleotides of the invention, but this is not essential. Most preferably, the genomic DNA is obtained from the biological material described herein in the Examples. Such libraries can be in vectors suitable for carrying large segments of a genome, such as PI or YAC, as described in detail in Sambrook et al., supra, 9.4-9.30. In addition, genomic sequences can be isolated from human BAC libraries, which are commercially available from Research Genetics, Inc., Huntsville, Alabama, USA, for example. In order to obtain additional 5° or 3° sequences, chromosome walking is performed, as described in Sambrook et al., such that adjacent and overlapping fragments of genomic DNA are isolated. These are mapped and pieced together, as is known in the art, using restriction divestion enzymes and DNA livase.

Using the polynucleotide sequences of the invention, corresponding full-length genes can be isolated using both classical and PCR methods to construct and probe cDNA libraries. Using either method, Northern blots, preferably, are performed on a number of cell types to determine which cell lines express the gene of interest at the highest level. Classical methods of constructing cDNA libraries are taught in Sambrook et al., supra. With these methods, cDNA can be produced from mRNA and inserted into viral or expression vectors. Typically, libraries of mRNA comprising poly(A) tails can be produced with poly(T) primers. Similarly, cDNA libraries can be produced using the instant sequences as primers.

PCR methods are used to amplify the members of a cDNA library that comprise the desired insert. In this case, the desired insert will contain sequence from the full length cDNA that corresponds to the instant polymucleotides. Such PCR methods include gene trapping and RACE methods. Gene trapping entails inserting a member of a cDNA library into a vector. The vector then is denatured to produce single stranded molecules. Next, a substrate-bound probe, such as a biotinylated oligo, is used to trap cDNA inserts of interest. Biotinylated probes can be linked to an avidin-bound solid substrate. PCR methods can be used to amplify the trapped cDNA. To trap sequences corresponding to the full length genes, the labeled probe sequence is based on the polynucleotide sequences of the invention. Random primers or primers specific to the library vector can be used to amplify the trapped cDNA. Such gene trapping techniques are described in Gruber et al., WO 95/04745 and Gruber et al., USPN 5,500,356. Kits are commercially available to perform gene trapping experiments from, for example, Life Technologies, Gaithersburg, Maryland, USA.

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"Rapid amplification of cDNA ends," or RACE, is a PCR method of amplifying cDNAs from a number of different RNAs. The cDNAs are ligated to an oligonucleotide linker, and amplified by PCR using two primers. One primer is based on sequence from the instant polynucleotides, for which full length sequence is desired, and a second primer comprises sequence that hybridizes to the oligonucleotide linker to amplify the cDNA. A description of this method is reported in WO 97/19110. In preferred embodiments of RACE, a common primer is designed to anneal to an arbitrary adaptor sequence ligated to cDNA ends (Apte and Siebert, Biotechniques (1993) 15.890-893; Edwards et al., Nuc. Acids Res. (1991) 19:5227-5232). When a single gene-specific RACE primer is paired with the common primer, preferential amplification of sequences between the single gene specific primer and the common primer occurs. Commercial cDNA pools modified for use in RACE are available.

Another PCR-based method generates full-length cDNA library with anchored ends without needing specific knowledge of the cDNA sequence. The method uses lock-docking primers (I-VI), where one primer, poly TV (I-III) locks over the polyA tail of eukaryotic mRNA producing first strand synthesis and a second primer, polyGH (IV-VI) locks onto the polyC tail added by terminal dooxynucleoidyl transferase (TdT)(see, e.g., WO 96/40998).

The promoter region of a gene generally is located 5° to the initiation site for RNA polymerase II. Hundreds of promoter regions contain the "TATA" box, a sequence such as TATTA or TATAA, which is sensitive to mutations. The promoter region can be obtained by performing 5° RACE using a primer from the coding region of the gene. Alternatively, the cDNA can be used as a probe for the genomic sequence, and the region 5° to the coding region is identified by "walking up." If the gene is

highly expressed or differentially expressed, the promoter from the gene can be of use in a regulatory construct for a heterologous gene.

Once the full-length cDNA or gene is obtained, DNA encoding variants can be prepared by site-directed mutagenesis, described in detail in Sambrook et al., 15.3-15.63. The choice of codon or nucleotide to be replaced can be based on disclosure herein on optional changes in amino acids to achieve altered protein structure and/or function.

As an alternative method to obtaining DNA or RNA from a biological material, nucleic acid comprising nucleotides having the sequence of one or more polynucleotides of the invention can be synthesized. Thus, the invention encompasses nucleic acid molecules ranging in length from 15 nt (corresponding to at least 15 contiguous nt of one of SEQ ID NOS:1-6010) up to a maximum length suitable for one or more biological manipulations, including replication and expression, of the nucleic acid molecule. The invention includes but is not limited to (a) nucleic acid having the size of a full gene, and comprising at least one of SEQ ID NOS:1-6010; (b) the nucleic acid of (a) also comprising at least one additional gene, operably linked to permit expression of a fusion protein; (c) an expression vector comprising (a) or (b); (d) a plasmid comprising (a) or (b); and (c) a recombinant viral particle comprising (a) or (b). Once provided with the polynucleotides disclosed herein, construction or preparation of (a) - (c) are well within the skill in the art.

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The sequence of a nucleic acid comprising at least 15 contiguous nt of at least any one of SEQ ID NOS:1-6010, preferably the entire sequence of at least any one of SEQ ID NOS:1-6010, is not limited and can be any sequence of A, T, G, and/or C (for DNA) and A, U, G, and/or C (for RNA) or modified bases thereof, including inosine and pseudouridine. The choice of sequence will depend on the desired function and can be dictated by coding regions desired, the intron-like regions desired, and the regulatory regions desired. Where the entire sequence of any one of SEQ ID NOS:1-6010 is within the nucleic acid, the nucleic acid obtained is referred to herein as a polynucleotide comprising the sequence of any one of SEO ID NOS:1-6010.

Expression of Polypeptide Encoded by Full-Length cDNA or Full-Length Gene

The provided polynucleotides (e.g., a polynucleotide having a sequence of one of SEQ ID NOS:1-6010), the corresponding cDNA, or the full-length gene is used to express a partial or complete gene product. Constructs of polynucleotides having sequences of SEQ ID NOS:1-6010 can also be generated synthetically. Alternatively, single-step assembly of a gene and entire plasmid from large numbers of oligodecoxyribonucleotides is described by, e.g., Stemmer et al., Gene (Amsterdam) (1995) 164(1):49-53. In this method, assembly PCR (the synthesis of long DNA sequences from large numbers of oligodecoxyribonucleotides (oligos)) is described. The method is derived from DNA

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shuffling (Stemmer, Nature (1994) 370:389-391), and does not rely on DNA ligase, but instead relies on DNA polymerase to build increasingly longer DNA fragments during the assembly process.

Appropriate polynucleotide constructs are purified using standard recombinant DNA techniques as described in, for example, Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd Ed., (1989) Cold Spring Harbor Press, Cold Spring Harbor, NY, and under current regulations described in United States Dept. of HHIS, National Institute of Health (NIH) Guidelines for Recombinant DNA Research. The gene product encoded by a polynucleotide of the invention is expressed in any expression system, including, for example, bacterial, yeast, insect, amphibian and mammalian systems. Vectors, host cells and methods for obtaining expression in same are well known in the art. Suitable vectors and host cells are described in USPN 5.654,173.

Polynucleotide molecules comprising a polynucleotide sequence provided herein are generally propagated by placing the molecule in a vector. Viral and non-viral vectors are used, including plasmids. The choice of plasmid will depend on the type of cell in which propagation is desired and the purpose of propagation. Certain vectors are useful for amplifying and making large amounts of the desired DNA sequence. Other vectors are suitable for expression in cells in culture. Still other vectors are suitable for transfer and expression in cells in a whole animal or person. The choice of appropriate vector is well within the skill of the art. Many such vectors are available commercially. Methods for preparation of vectors comprising a desired sequence are well known in the art.

The polynucleotides set forth in SEQ ID NOS:1-6010 or their corresponding full-length polynucleotides are linked to regulatory sequences as appropriate to obtain the desired expression properties. These can include promoters (attached either at the 5' end of the sense strand or at the 3' end of the antisense strand), enhancers, terminators, operators, repressors, and inducers. The promoters can be regulated or constitutive. In some situations it may be desirable to use conditionally active promoters, such as tissue-specific or developmental stage-specific promoters. These are linked to the desired nucleotide sequence using the techniques described above for linkage to vectors. Any techniques known in the art can be used.

When any of the above host cells, or other appropriate host cells or organisms, are used to replicate and/or express the polynucleotides or nucleic acids of the invention, the resulting replicated nucleic acid, RNA, expressed protein or polypeptide, is within the scope of the invention as a product of the host cell or organism. The product is recovered by any appropriate means known in the art.

Once the gene corresponding to a selected polymucleotide is identified, its expression can be regulated in the cell to which the gene is native. For example, an endogenous gene of a cell can be regulated by an exogenous regulatory sequence as disclosed in USPN 5.641.670.

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Identification of Functional and Structural Motifs

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Translations of the nucleotide sequence of the provided polynucleotides, cDNAs or full genes can be aligned with individual known sequences. Similarity with individual sequences can be used to determine the activity of the polypeptides encoded by the polynucleotides of the invention. Also, sequences exhibiting similarity with more than one individual sequence can exhibit activities that are characteristic of either or both individual sequences.

The full length sequences and fragments of the polynucleotide sequences of the nearest neighbors can be used as probes and primers to identify and isolate the full length sequence corresponding to provided polynucleotides. The nearest neighbors can indicate a tissue or cell type to be used to construct a library for the full-length sequences corresponding to the provided polynucleotides.

Typically, a selected polynucleotide is translated in all six frames to determine the best alignment with the individual sequences. The sequences disclosed herein in the Sequence Listing are in a 5' to 3' orientation and translation in three frames can be sufficient (with a few specific exceptions as described in the Examples). These amino acid sequences are referred to, generally, as query sequences, which will be aligned with the individual sequences. Databases with individual sequences are described in "Computer Methods for Macromolecular Sequence Analysis" Methods in Enzymology (1996) 266, Doolittle, Academic Press, Inc., a division of Harcourt Brace & Co., San Diego, California, USA. Databases include GenBank, EMBL, and DNA Database of Japan (DDBJ).

Query and individual sequences can be aligned using the methods and computer programs described above, and include BLAST 2.0, available over the world wide web at a site supported by the National Center for Biotechnology Information, which is supported by the National Library of Medicine and the National Institutes of Health. See also Altschul, et al. Nucleic Acids Res. (1997) 25:3389-3402. Another alignment algorithm is Fasta, available in the Genetics Computing Group (GCG) package, Madison, Wisconsin, USA, a wholly owned subsidiary of Oxford Molecular Group, Inc. Other techniques for alignment are described in Doolittle, supra. Preferably, an alignment program that permits gaps in the sequence is utilized to align the sequences. The Smith-Waterman is one type of algorithm that permits gaps in sequence alignments. See Meth. Mol. Biol. (1997) 70: 173-187. Also, the GAP program using the Necdleman and Wunsch alignment method can be utilized to align sequences. An alternative search strategy uses MPSRCH software, which runs on a MASPAR computer. MPSRCH uses a Smith-Waterman algorithm to score sequences on a massively parallel computer. This approach improves ability to identify sequences that are distantly related matches, and is especially tolerant of small gaps and mucleotide sequence errors. Amino acid sequences encoded by the provided polynucleotides can be used to search both protein and DNA databases. Incorporated

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herein by reference are all sequences that have been made public as of the filing date of this application by any of the DNA or protein sequence databases, including the patent databases (e.g., GeneSeq). Also incorporated by reference are those sequences that have been submitted to these databases as of the filing date of the present application but not made public until after the filing date of the present application.

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Results of individual and query sequence alignments can be divided into three categories: high similarity, weak similarity, and no similarity. Individual alignment results ranging from high similarity to weak similarity around a basis for determining polypeptide activity and/or structure. Parameters for categorizing individual results include: percentage of the alignment region length where the strongest alignment is found, percent sequence identity, and p value. The percentage of the alignment region length is calculated by counting the number of residues of the individual sequence found in the region of strongest alignment, e.g., contiguous region of the individual sequence that contains the greatest number of residues that are identical to the residues of the corresponding region of the aligned query sequence. This number is divided by the total residue length of the query sequence to calculate a percentage. For example, a query sequence C20 amino acid residues might be aligned with a 20 amino acid region of an individual sequence. The individual sequence might be identical to amino acid residues 5, 9-15, and 17-19 of the query sequence. The region of strongest alignment is thus the region stretching from residue 9-19, an 11 amino acid stretch. The percentage of the alignment region length is: 11 (length of the region of strongest alignment) divided by (query sequence length) 20 or 55%.

Percent sequence identity is calculated by counting the number of amino acid matches between the query and individual sequence and dividing total number of matches by the number of residues of the individual sequences found in the region of strongest alignment. Thus, the percent identity in the example above would be 10 matches divided by 11 amino acids, or approximately, 90.9%

P value is the probability that the alignment was produced by chance. For a single alignment, the p value can be calculated according to Karlin et al., Proc. Natl. Acad. Sci. (1990) 87:2264 and Karlin et al., Proc. Natl. Acad. Sci. (1993) 90. The p value of multiple alignments using the same query sequence can be calculated using an heuristic approach described in Altschul et al., Nat. Genet. (1994) 6:119. Alignment programs such as BLAST program can calculate the p value. See also Altschul et al., Nucleic Acids Res. (1997) 25:3389-3402.

Another factor to consider for determining identity or similarity is the location of the similarity or identity. Strong local alignment can indicate similarity even if the length of alignment is short. Sequence identity scattered throughout the length of the query sequence also can indicate a similarity between the query and profile sequences. The boundaries of the region where the sequences align can be determined according to Doolittle, supra; BIAST 2.0 (see, e.g., Altschul, et al. Nucleic Acids Res.

(1997) 25:3389-3402) or FAST programs; or by determining the area where sequence identity is highest.

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High Similarity. In general, in alignment results considered to be of high similarity, the percent of the alignment region length is typically at least about 55% of total length query sequence; more typically, at least about 58%; even more typically; at least about 60% of the total residue length of the alignment region can be as much as about 62%; more usually, as much as about 64%; even more usually, as much as about 66%. Further, for high similarity, the region of alignment, typically, exhibits at least about 75% of sequence identity; more typically, at least about 78%; even more typically; at least about 80% sequence identity. Usually, percent sequence identity can be as much as about 82%; more usually, as much as about 84%; even more usually, as much as about 84%; even more usually, as much as about 84%;

The p value is used in conjunction with these methods. If high similarity is found, the query sequence is considered to have high similarity with a profile sequence when the p value is less than or equal to about 10e-2; more usually; less than or equal to about 10e-3; even more usually; less than or equal to about 10e-4. More typically, the p value is no more than about 10e-5; more typically, no more than or equal to about 10e-10; even more typically, no more than or equal to about 10e-15 for the query sequence to be considered high similarity.

Weak Similarity. In general, where alignment results considered to be of weak similarity, there is no minimum percent length of the alignment region nor minimum length of alignment. A better showing of weak similarity is considered when the region of alignment is, typically, at least about 15 amino acid residues in length; more typically, at least about 20; even more typically, at least about 25 amino acid residues in length. Usually, length of the alignment region can be as much as about 30 amino acid residues; more usually, as much as about 40; even more usually, as much as about 40 amino acid residues. Further, for weak similarity, the region of alignment, typically, exhibits at least about 35% of sequence identity; more typically, at least about 40%, even more typically, at least about 45% sequence identity. Usually, percent sequence identity can be as much as about 50%; more usually, as much as about 55%; even more usually, as much as about 50%;

If low similarity is found, the query sequence is considered to have weak similarity with a profile sequence when the p value is usually less than or equal to about 10e-2; more usually, less than or equal to about 10e-3. Wore typically, the p value is no more than about 10e-5; more usually; no more than or equal to about 10e-10; even more usually, no more than or equal to about 10e-10; even more usually, no more than or equal to about 10e-10; and to about 10e-10 for the query sequence to be considered weak similarity.

Similarity Determined by Sequence Identity Alone. Sequence identity alone can be used to determine similarity of a query sequence to an individual sequence and can indicate the activity of the

sequence. Such an alignment, preferably, permits gaps to align sequences. Typically, the query sequence is related to the profile sequence if the sequence identity over the entire query sequence is at least about 15%; more typically, at least about 25%; even more typically, at least about 55%. Sequence identity alone as a measure of similarity is most useful when the query sequence is usually, at least 80 residues in length; more usually, at least 90 residues in length; when the query sequence is usually, at least 95 amino acid residues in length. More typically, similarity can be concluded based on sequence identity alone when the query sequence is preferably 100 residues in length; more preferably, 120 residues in length; over more preferably, 150 amino acid residues in length;

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Alignments with Profile and Multiple Aligned Sequences. Translations of the provided polynucleotides can be aligned with amino acid profiles that define either protein families or common motifs. Also, translations of the provided polynucleotides can be aligned to multiple sequence alignments (MSA) comprising the polypeptide sequences of members of protein families or motifs. Similarity or identity with profile sequences or MSAs can be used to determine the activity of the gene products (e.g., polypeptides) encoded by the provided polynucleotides or corresponding cDNA or genes. For example, sequences that show an identity or similarity with a chemokine profile or MSA can exhibit chemokine activities.

Profiles can be designed manually by (1) creating an MSA, which is an alignment of the amino acid sequence of members that belong to the family and (2) constructing a statistical representation of the alignment. Such methods are described, for example, in Birney et al., Nucl. Acid Res. (1996) 424(14): 2730-2739. MSAs of some protein families and motifs are publicly available. For example, the Genome Sequencing Center at thw Washington University School of Medicine provides a web set (Pfam) which provides MSAs of 547 different families and motifs. These MSAs are described also in Sonnhammer et al., Proteins (1997) 28: 405-420. Other sources over the world wide web include the site supported by the European Molecular Biology Laboratories in Heidelberg, Germany. A brief description of these MSAs is reported in Pascarella et al., Prot. Eng. (1996) 9(3):249-251. Techniques for building profiles from MSAs are described in Sonnhammer et al., supra; Birney et al., supra; and "Computer Methods for Macromolecular Sequence Analysis," Methods in Enzymology (1996) 266, Doolittle, Academie Press, Inc., San Diego, California, USA.

Similarity between a query sequence and a protein family or motif can be determined by (a) comparing the query sequence against the profile and/or (b) aligning the query sequence with the members of the family or motif. Typically, a program such as Searchwise is used to compare the query sequence to the statistical representation of the multiple alignment, also known as a profile (see Birney

et al., supra). Other techniques to compare the sequence and profile are described in Sonnhammer et al., supra and Doolittle, supra.

Next, methods described by Feng et al., J. Mol. Evol. (1987) 25:351 and Higgins et al.,

CABIOS (1989) 5:151 can be used align the query sequence with the members of a family or motif,
also known as a MSA. Sequence alignments can be generated using any of a variety of software tools.

Examples include PileUp, which creates a multiple sequence alignment, and is described in Feng et al.,
J. Mol. Evol. (1987) 25:351. Another method, GAP, uses the alignment method of Needleman et al., J.

Mol. Biol. (1970) 48:443. GAP is best suited for global alignment of sequences. A third method,
BestFit, functions by inserting gaps to maximize the number of matches using the local homology
algorithm of Smith et al., Adv. Appl. Math. (1981) 2:482. In general, the following factors are used to
determine if a similarity between a query sequence and a profile or MSA exists: (1) number of
conserved residues found in the query sequence, (2) percentage of conserved residues found in the
query sequence. (3) number of frameshifts, and (4) spacins between conserved residues.

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Some alignment programs that both translate and align sequences can make any number of frameshifts when translating the nucleotide sequence to produce the best alignment. The fewer frameshifts needed to produce an alignment, the stronger the similarity or identity between the query and profile or MSAs. For example, a weak similarity resulting from no frameshifts can be a better indication of activity or structure of a query sequence, than a strong similarity resulting from two frameshifts. Preferably, three or fewer frameshifts are found in an alignment; more preferably two or fewer frameshifts; even more preferably, no frameshifts are found in an alignment of query and profile or MSAs.

Conserved residues are those amino acids found at a particular position in all or some of the family or motif members. Alternatively, a position is considered conserved if only a certain class of amino acids is found in a particular position in all or some of the family members. For example, the N-terminal position can contain a positively charged amino acid, such as lysine, arginine, or histidine,

Typically, a residue of a polypeptide is conserved when a class of amino acids or a single amino acid is found at a particular position in at least about 40% of all class members; more typically, at least about 50%, even more typically, at least about 60% of the members. Usually, a residue is conserved when a class or single amino acid is found in at least about 70% of the members of a family or motif; more usually, at least about 80%; even more usually, at least about 90%; even more usually, at least about 95%.

A residue is considered conserved when three unrelated amino acids are found at a particular position in some or all of the members; more usually, two unrelated amino acids. These residues are conserved when the unrelated amino acids are found at particular positions in at least about 40% of all

class member, more typically, at least about 50%; even more typically, at least about 60% of the members. Usually, a residue is conserved when a class or single amino acid is found in at least about 70% of the members of a family or motif; more usually, at least about 80%; even more usually, at least about 90%; even more usually, at least about 90%; even more usually, at least about 90%.

A query sequence has similarity to a profile or MSA when the query sequence comprises at least about 25% of the conserved residues of the profile or MSA; more usually, at least about 30%; even more usually; at least about 40%. Typically, the query sequence has a stronger similarity to a profile sequence or MSA when the query sequence comprises at least about 45% of the conserved residues of the profile or MSA; more typically, at least about 50%; even more typically, at least about 55%.

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Identification of Secreted & Membrane-Bound Polypeptides. Both secreted and membrane-bound polypeptides of the present invention are of particular interest. For example, levels of secreted polypeptides can be assayed in body fluids that are convenient, such as blood, plasma, serum, and other body fluids such as urine, prostatic fluid and semen. Membrane-bound polypeptides are useful for constructing vaccine antigens or inducing an inmune response. Such antigens would comprise all or part of the extracellular region of the membrane-bound polypeptides. Because both secreted and membrane-bound polypeptides comprise a fragment of contiguous hydrophobic amino acids, hydrophobicity predicting algorithms can be used to identify such polypeptides.

A signal sequence is usually encoded by both secreted and membrane-bound polypeptide genes to direct a polypeptide to the surface of the cell. The signal sequence usually comprises a stretch of hydrophobic residues. Such signal sequences can fold into helical structures. Membrane-bound polypeptides typically comprise at least one transmembrane region that possesses a stretch of hydrophobic amino acids that can transverse the membrane. Some transmembrane regions also exhibit a helical structure. Hydrophobic fragments within a polypeptide can be identified by using computer algorithms. Such algorithms include Hopp & Woods, Proc. Natl. Acad. Sci. USA (1981) 78:3824-3828; Kyte & Doolittle, J. Mol. Biol. (1982) 157: 105-132; and RAOAR algorithm, Degli Esposti et al., Eur. J. Biochem. (1990) 190: 207-219.

Another method of identifying secreted and membrane-bound polypeptides is to translate the polynucleotides of the invention in all six frames and determine if at least 8 contiguous hydrophobic amino acids are present. Those translated polypeptides with at least 8; more typically, 10; even more typically, 12 contiguous hydrophobic amino acids are considered to be either a putative secreted or membrane bound polypeptide. Hydrophobic amino acids include alanine, glycine, histidine, isoleucine, leucine, lysine, methionine, phenylalanine, proline, threonine, tryptophan, tyrosine, and valine

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Identification of the Function of an Expression Product of a Full-Length Gene

Ribozymes, antisense constructs, and dominant negative mutants can be used to determine function of the expression product of a gene corresponding to a polynucleotide provided herein. These methods and compositions are particularly useful where the provided novel polynucleotide exhibits no significant or substantial homology to a sequence encoding a gene of known function. Antisense molecules and ribozymes can be constructed from synthetic polynucleotides. Typically, the phosphoramidite method of oligonucleotide synthesis is used. See Beaucage et al., Tet. Lett. (1981) 22:1859 and USPN 4,668,777. Automated devices for synthesis are available to create oligonucleotides using this chemistry. Examples of such devices include Biosearch 8600, Models 392 and 394 by Applied Biosystems, a division of Perkin-Elmer Corp., Foster City, California, USA; and Expedite by Perceptive Biosystems, Framingham, Massachusetts, USA. Synthetic RNA, phosphate analog oligonucleotides, and chemically derivatized oligonucleotides can also be produced, and can be covalently attached to other molecules. RNA oligonucleotides can be synthesized, for example, using RNA phosphoramidites. This method can be performed on an automated synthesizer, such as Applied Biosystems, Models 392 and 394, Foster City, California, USA.

Phosphorothioate oligonucleotides can also be synthesized for antisense construction. A sulfurizing reagent, such as tetracthythirnam disulfide (TETD) in acetonitrile can be used to convert the internucleotide cyanocthyl phosphite to the phosphorothioate triester within 15 minutes at room temperature. TETD replaces the iodine reagent, while all other reagents used for standard phosphoramidite chemistry remain the same. Such a synthesis method can be automated using Models 392 and 394 by Applied Biosystems, for example.

Oligonucleotides of up to 200 nt can be synthesized, more typically, 100 nt; more typically 50 nt; even more typically, 30 to 40 nt. These synthetic fragments can be annealed and ligated together to construct larger fragments. See, for example, Sambrook et al., supra. Trans-cleaving catalytic RNAs (ribozymes) are RNA molecules possessing endoribonuclease activity. Ribozymes are specifically designed for a particular target, and the target message must contain a specific nucleotide sequence. They are engineered to cleave any RNA species site-specifically in the background of cellular RNA. The cleavage event renders the mRNA unstable and prevents protein expression. Importantly, ribozymes can be used to inhibit expression of a gene of unknown function for the purpose of determining its function in an in vitro or in vivo context, by detecting the phenotypic effect. One commonly used ribozyme motif is the hammerhead, for which the substrate sequence requirements are minimal. Design of the hammerhead ribozyme, as well as therapeutic uses of ribozymes, are disclosed in Usman et al., Current Opin. Struct. Biol. (1996) 6:527. Methods for production of ribozymes,

including hairpin structure ribozyme fragments, methods of increasing ribozyme specificity, and the like are known in the art.

The hybridizing region of the ribozyme can be modified or can be prepared as a branched structure as described in Horn and Urdea, Nucleic Acids Res. (1989) 17:6959. The basic structure of the ribozymes can also be chemically altered in ways familiar to those skilled in the art, and chemically synthesized ribozymes can be administered as synthetic oligonucleotide derivatives modified by monomeric units. In a therapeutic context, liposome mediated delivery of ribozymes improves cellular uptake, as described in Birikh et al., Eur. J. Biochem. (1997) 245:1.

Antisense nucleic acids are designed to specifically bind to RNA, resulting in the formation of RNA-DNA or RNA-RNA hybrids, with an arrest of DNA replication, reverse transcription or messenger RNA translation. Antisense polymucleotides based on a selected polymucleotide sequence can interfere with expression of the corresponding gene. Antisense polymucleotides are typically generated within the cell by expression from antisense constructs that contain the antisense strand as the transcribed strand. Antisense polymucleotides based on the disclosed polymucleotides will bind and/or interfere with the translation of mRNA comprising a sequence complementary to the antisense polymucleotide. The expression products of control cells and cells treated with the antisense construct are compared to detect the protein product of the gene corresponding to the polymucleotide upon which the antisense construct is based. The protein is isolated and identified using routine biochemical methods.

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Given the extensive background literature and clinical experience in antisense therapy, one skilled in the art can use selected polynucleotides of the invention as additional potential therapeutics. The choice of polynucleotide can be narrowed by first testing them for binding to "hot spot" regions of the genome of cancerous cells. If a polynucleotide is identified as binding to a "hot spot," testing the polynucleotide as an antisense compound in the corresponding cancer cells is warranted.

As an alternative method for identifying function of the gene corresponding to a polynucleotide disclosed herein, dominant negative mutations are readily generated for corresponding proteins that are active as homomultimers. A mutant polypeptide will interact with wild-type polypeptides (made from the other allele) and form a non-functional multimer. Thus, a mutation is in a substrate-binding domain, a catalytic domain, or a cellular localization domain. Preferably, the mutant polypeptide will be overproduced. Point mutations are made that have such an effect. In addition, fusion of different polypeptides of various lengths to the terminus of a protein can yield dominant negative mutants. General strategies are available for making dominant negative mutants (sec, e.g., Herskowitz, Nature (1987) 329-219). Such techniques can be used to create loss of function mutations, which are useful for determining protein function.

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Polypeptides and Variants Thereof

The polypeptides of the invention include those encoded by the disclosed polymuclectides, as well as nucleic acids that, by virtue of the degeneracy of the genetic code, are not identical in sequence to the disclosed polynuclectides. Thus, the invention includes within its scope a polypeptide encoded by a polynuclectide having the sequence of any one of SEQ ID NOS:1-6010 or a variant thereof.

In general, the term "polypeptide" as used herein refers to both the full length polypeptide encoded by the recited polynucleotide, the polypeptide encoded by the gene represented by the recited polynucleotide, as well as portions or fragments thereof. "Polypeptides" also includes variants of the naturally occurring proteins, where such variants are homologous or substantially similar to the naturally occurring protein, and can be of an origin of the same or different species as the naturally occurring protein (e.g., human, murine, or some other species that naturally expresses the recited polypeptide, usually a mammalian species). In general, variant polypeptides have a sequence that has at least about 80%, usually at least about 90%, and more usually at least about 98% sequence identity with a differentially expressed polypeptide of the invention, as measured by BLAST 2.0 using the parameters described above. The variant polypeptides can be naturally or non-naturally glycosylated, i.e., the polypeptide has a glycosylation pattern that differs from the glycosylation pattern found in the corresponding naturally occurring protein.

The invention also encompasses homologs of the disclosed polypeptides (or fragments thereof) where the homologs are isolated from other species, i.e. other animal or plant species, where such homologs, usually mammalian species, e.g. rodents, such as mice, rats; domestic animals, e.g., horse, cow, dog, cat; and humans. By "homologs" is meant a polypeptide having at least about 35%, usually at least about 40% and more usually at least about 60% amino acid sequence identity to a particular differentially expressed protein as identified above, where sequence identity is determined using the BLAST 2.0 algorithm, with the parameters described surra.

In general, the polypeptides of the subject invention are provided in a non-naturally occurring environment, e.g. are separated from their naturally occurring environment. In certain embodiments, the subject protein is present in a composition that is enriched for the protein as compared to a control. As such, purified polypeptide is provided, where by purified is meant that the protein is present in a composition that is substantially free of non-differentially expressed polypeptides, where by substantially free is meant that less than 90%, usually less than 60% and more usually less than 50% of the composition is made up of non-differentially expressed polypeptides.

Also within the scope of the invention are variants; variants of polypeptides include mutants, fragments, and fusions. Mutants can include amino acid substitutions, additions or deletions. The amino acid substitutions can be conservative amino acid substitutions or substitutions to eliminate non-

essential amino acids, such as to alter a glycosylation site, a phosphorylation site or an acetylation site. or to minimize misfolding by substitution or deletion of one or more cysteine residues that are not necessary for function. Conservative amino acid substitutions are those that preserve the general charge. hydrophobicity/ hydrophilicity, and/or steric bulk of the amino acid substituted. Variants can be designed so as to retain or have enhanced biological activity of a particular region of the protein (e.g., a functional domain and/or, where the polypeptide is a member of a protein family, a region associated with a consensus sequence). Selection of amino acid alterations for production of variants can be based upon the accessibility (interior vs. exterior) of the amino acid (see, e.g., Go et al. Int. J. Peptide Protein Res. (1980) 15:211), the thermostability of the variant polypeptide (see, e.g., Querol et al., Prot. Eng. (1996) 9:265), desired glycosylation sites (see, e.g., Olsen and Thomsen, J. Gen. Microbiol. (1991) 137:579), desired disulfide bridges (sec, e.g., Clarke et al., Biochemistry (1993) 32:4322; and Wakarchuk et al., Protein Eng. (1994) 7:1379), desired metal binding sites (see, e.g., Toma et al., Biochemistry (1991) 30:97, and Haezerbrouck et al., Protein Eng. (1993) 6:643), and desired substitutions within proline loops (see, e.g., Masul et al., Appl. Env. Microbiol. (1994) 60:3579). Cysteine-depleted muteins can be produced as disclosed in USPN 4,959,314. Variants also include fragments of the polypeptides disclosed herein, particularly haptens, biologically active fragments, and/or fragments corresponding to functional domains. Fragments of

Variants also include fragments of the polypeptides disclosed herein, particularly haptens, biologically active fragments, and/or fragments corresponding to functional domains. Fragments of interest will typically be at least about 10 as to at least about 15 as in length, usually at least about 50 as in length, and can be as long as 300 as in length or longer, but will usually not exceed about 1000 as in length, where the fragment will have a stretch of amino acids that is identical to a polypeptide encoded by a polynucleotide having a sequence of any SEQ ID NOS:1-6010, or a homolog thereof. The protein variants described herein are encoded by polynucleotides that are within the scope of the invention. The genetic code can be used to select the appropriate codons to construct the corresponding variants.

25 Computer-Related Embodiments

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In general, a library of polynucleotides is a collection of sequence information, which information is provided in either biochemical form (e.g., as a collection of polynucleotide molecules), or in electronic form (e.g., as a collection of polynucleotide sequences stored in a computer-readable form, as in a computer system and/or as part of a computer program). The sequence information of the polynucleotides can be used in a variety of ways, e.g., as a resource for gene discovery, as a representation of sequences expressed in a selected cell type (e.g., cell type markers), and/or as markers of a given disease or disease state. In general, a disease marker is a representation of a gene product that is present in all cells affected by disease cither at an increased or decreased level relative to a normal cell (e.g., a cell of the same or similar type that is not substantially affected by disease). For

example, a polynucleotide sequence in a library can be a polynucleotide that represents an mRNA, polypeptide, or other gene product encoded by the polynucleotide, that is either overexpressed or underexpressed in a breast ductal cell affected by cancer relative to a normal (i.e., substantially diseasefree) breast cell.

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The nucleotide sequence information of the library can be embodied in any suitable form, e.g., electronic or biochemical forms. For example, a library of sequence information embodied in electronic form comprises an accessible computer data file (or, in biochemical form, a collection of nucleic acid molecules) that contains the representative nucleotide sequences of genes that are differentially expressed (e.g., overexpressed or underexpressed) as between, for example, i) a cancerous cell and a normal cell; ii) a cancerous cell and a dysplastic cell; iii) a cancerous cell and a cell affected by a disease or condition other than cancer; iv) a metastatic cancerous cell and a normal cell and/or non-metastatic cancerous cell; v) a malignant cancerous cell and a non-malignant cancerous cell (or a normal cell) and/or vi) a dysplastic cell relative to a normal cell. Other combinations and comparisons of cells affected by various diseases or stages of disease will be readily apparent to the ordinarily skilled artisan. Biochemical embodiments of the library include a collection of nucleic acids that have the sequences of the genes in the library, where the nucleic acids can correspond to the entire gene in the library or to a fragment thereof, as described in greater detail below.

The polynucleotide libraries of the subject invention generally comprise sequence information of a purality of polynucleotide sequences, where at least one of the polynucleotides has a sequence of any of SEQ ID NOS:1-6010. By plurality is meant at least 2, usually at least 3 and can include up to all of SEQ ID NOS:1-6010. The length and number of polynucleotides in the library will vary with the nature of the library, e.g., if the library is an oligonucleotide array, a cDNA array, a computer database of the sequence information, etc.

Where the library is an electronic library, the nucleic acid sequence information can be present in a variety of media. "Media" refers to a manufacture, other than an isolated nucleic acid molecule, that contains the sequence information of the present invention. Such a manufacture provides the genome sequence or a subset thereof in a form that can be examined by means not directly applicable to the sequence as it exists in a nucleic acid. For example, the nucleotide sequence of the present invention, e.g. the nucleic acid sequences of any of the polynucleotides of SEQ ID NOS:1-6010, can be recorded on computer readable media, e.g. any medium that can be read and accessed directly by a computer. Such media include, but are not limited to: magnetic storage media, such as a floppy disc, a hard disc storage medium, and a magnetic tape; optical storage media such as CD-ROM; electrical storage media such as RAM and ROM; and hybrids of these categories such as magnetic/optical storage media. One of skill in the art can readily appreciate how any of the presently known computer

readable mediums can be used to create a manufacture comprising a recording of the present sequence information. "Recorded" refers to a process for storing information on computer readable medium, using any such methods as known in the art. Any convenient data storage structure can be chosen, based on the means used to access the stored information. A variety of data processor programs and formats can be used for storage, e.g. word processing text file, database format, etc. In addition to the sequence information, electronic versions of the libraries of the invention can be provided in conjunction or connection with other computer-readable information and/or other types of computer-readable files (e.g., searchable files, executable files, etc. including, but not limited to, for example, search process means of the search process of t

By providing the nucleotide sequence in computer readable form, the information can be accessed for a variety of purposes. Computer software to access sequence information is publicly available. For example, the gapped BLAST (Altschul et al. Nucleic Acids Res. (1997) 25:3389-3402) and BLAZE (Brutlag et al. Comp. Chem. (1993) 17:203) search algorithms on a Sybase system can be used to identify open reading frames (ORFs) within the genome that contain homology to ORFs from other organisms.

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As used herein, "a computer-based system" refers to the hardware means, software means, and data storage means used to analyze the nucleotide sequence information of the present invention. The minimum hardware of the computer-based systems of the present invention comprises a central processing unit (CPU), input means, output means, and data storage means. A skilled artisan can readily appreciate that any one of the currently available computer-based system are suitable for use in the present invention. The data storage means can comprise any manufacture comprising a recording of the present sequence information as described above, or a memory access means that can access such a manufacture.

"Search means" refers to one or more programs implemented on the computer-based system, to compare a target sequence or target structural motif, or expression levels of a polynucleotide in a sample, with the stored sequence information. Search means can be used to identify fragments or regions of the genome that match a particular target sequence or target motif. A variety of known algorithms are publicly known and commercially available, e.g. MacPattern (EMBL), BLASTN and BLASTX (NCBI). A "target sequence" can be any polynucleotide or amino acid sequence of six or more contiguous nucleotides or two or more amino acids, preferably from about 10 to 100 amino acids or from about 30 to 300 nt A variety of comparing means can be used to accomplish comparison of sequence information from a sample (e.g., to analyze target sequences, target motifs, or relative expression levels) with the data storage means. A skilled artisan can readily recognize that any one of the publicly available homology search programs can be used as the search means for the computer

based systems of the present invention to accomplish comparison of target sequences and motifs.

Computer programs to analyze expression levels in a sample and in controls are also known in the art.

A "target structural motif," or "target motif," refers to any rationally selected sequence or combination of sequences in which the sequence(s) are chosen based on a three-dimensional configuration that is formed upon the folding of the target motif, or on consensus sequences of regulatory or active sites. There are a variety of target motifs known in the art. Protein target motifs include, but are not limited to, enzyme active sites and signal sequences. Nucleic acid target motifs include, but are not limited to, hairpin structures, promoter sequences and other expression elements such as binding sites for transcription factors.

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A variety of structural formats for the input and output means can be used to input and output the information in the computer-based systems of the present invention. One format for an output means ranks the relative expression levels of different polynucleotides. Such presentation provides a skilled artisan with a ranking of relative expression levels to determine a gene expression profile.

As discussed above, the "library" of the invention also encompasses biochemical libraries of the polynucleotides of SEQ ID NOS:1-6010, e.g., collections of nucleic acids representing the provided polynucleotides. The biochemical libraries can take a variety of forms, e.g., a solution of cDNAs, a pattern of probe nucleic acids stably associated with a surface of a solid support (i.e., an array) and the like. Of particular interest are nucleic acid arrays in which one or more of SEQ ID NOS:1-6010 is represented on the array. By array is meant an arricle of manufacture that has at least a substrate with at least two distinct nucleic acid targets on one of its surfaces, where the number of distinct nucleic acids can be considerably higher, typically being at least 10, usually at least 20, and often at least 25 distinct nucleic acid molecules. A variety of different array formats have been developed and are known to those of skill in the art. The arrays of the subject invention find use in a variety of applications, including gene expression analysis, drug screening, mutation analysis and the like, as disclosed in the above-listed exemplary patent documents.

In addition to the above nucleic acid libraries, analogous libraries of polypeptides are also provided, where the polypeptides of the library will represent at least a portion of the polypeptides encoded by a gene corresponding to one or more of SEQ ID NOS:1-6010. Utilities

The polynucleotides of the invention are useful in a variety of applications. Exemplary utilies of the polynucleotides of the invention are described below.

Construction of Larger Molecules: Recombinant DNAs and Nucleic Acid Multimers. In one embodiment of particular interest, the polynucleotides described herein as useful as the building blocks for larger molecules. In one example, the polynucleotide is a component of a larger cDNA molecule

which in turn can be adapted for expression in a host cell (e.g., a bacterial or eukaryotic (e.g., yeast or mammalian) host cell). The cDNA can include, in addition to the polypeptide encoded by the starting material polynucleotide (i.e., a polynucleotide described herein), an amino acid sequence that is heterologous to the polypeptide encoded by the polynucleotide described herein (e.g., as in a sequence encoding a fusion protein). In some embodiments, the polynucleotides described secriber in sused as starting material polynucleotide for synthesizing all or a portion of the gene to which the described polynucleotide corresponds. For example, a DNA molecule encoding a full-length human polypeptide can be constructed using a polynucleotide described herein as starting material.

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In another embodiment, the polynucleotides of the invention are used in nucleic acid multimers . Nucleic acid multimers can be linear or branched polymers of the same repeating single-stranded oligonucleotide unit or different single-stranded oligonucleotide units. Where the molecules are branched, the multimers are generally described as either "fork" or "comb" structures. The oligonucleotide units of the multimer may be composed of RNA, DNA, modified nucleotides or combinations thereof. At least one of the units has a sequence, length, and composition that permits it to bind specifically to a first single-stranded nucleotide sequence of interest, typically analyte or an oligonucleotide bound to the analyte. In order to achieve such specificity and stability, this unit will normally be 15 to 50 nt, preferably 15 to 30 nt, in length and have a GC content in the range of 40% to 60%. In addition to such unit(s), the multimer includes a multiplicity of units that are capable of hybridizing specifically and stably to a second single-stranded nucleotide of interest, typically a labeled oligonucleotide or another multimer. These units will also normally be 15 to 50 nt, preferably 15 to 30 nt, in length and have a GC content in the range of 40% to 60%. When a multimer is designed to be hybridized to another multimer, the first and second oligonucleotide units are heterogeneous (different). One or more of the polynucleotides described herein, or a portion of a polynucleotide described herein. can be used as a repeating unit of such nucleic acid multimers.

The total number of oligonucleotide units in the multimer will usually be in the range of 3 to 50, more usually 10 to 20. In multimers in which the unit that hybridizes to the nucleotide sequence of interest is different from the unit that hybridizes to the labeled oligonucleotide, the number ratio of the latter to the former will usually be 2:1 to 30:1, more usually 5:1 to 20:1, and-oreferably 10:1 to 15:1.

The oligonucleotide units of the multimer may be covalently linked directly to each other through phosphodiester bonds or through interposed linking agents such as mucleic acid, amino acid, carbohydrate or polyol bridges, or through other cross-linking agents that are capable of cross-linking nucleic acid or modified nucleic acid strands. The site(s) of linkage may be at the ends of the unit (in either normal 3,-5' orientation or randomly oriented) and/or at one or more internal nucleotides in the strand. In linear multimers the individual units are linked end-to-end to form a linear polymer. In one

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type of branched multimer three or more oligonucleotide units emanate from a point of origin to form a branched structure. The point of origin may be another oligonucleotide unit or a multifunctional molecule to which at least three units can be covalently bound. In another type, there is an oligonucleotide unit backbone with one or more pendant oligonucleotide units. These latter-type multimers are "fork-like", "comb-like" or combination "fork-" and "comb-like" in structure. The pendant units will normally depend from a modified nucleotide or other organic moiety having appropriate functional groups to which oligonucleotides may be conjugated or otherwise attached. The multimer may be totally linear, totally branched, or a combination of linear and branched portions. Preferably there will be at least two branch points in the multimer, more preferably at least 3, preferably 5 to 10. The multimer may include one or more segments of double-stranded sequences.

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Multimeric nucleic acid molecules are useful in amplifying the signal that results from hybridization of one the first sequence of the multimeric molecule to a target sequence. The amplification is theoretically proportional to the number of iterations of the second segment.

Without being held to theory, forked structures of greater than about eight branches exhibited steric hindrance which inhibited binding of labeled probes to the multimer. On the other hand, comb structures exhibit little or no steric problems and are thus a preferred type of branched multimer. For a description of branched nucleic acid multimers of both the fork and comb types, as well as methods of use and synthesis, see, e.g., U.S. Pat. Nos. 5,124,246 (fork-type structures); 5,710,264 (synthesis of comb structures); and 5,849,481.

Use of Polynucleotide Probes in Mapping, and in Tissue Profiling. Polynucleotide probes, generally comprising at least 12 contiguous nt of a polynucleotide as shown in the Sequence Listing, are used for a variety of purposes, such as chromosome mapping of the polynucleotide and detection of transcription levels. Additional disclosure about preferred regions of the disclosed polynucleotide sequences is found in the Examples. A probe that hybridizes specifically to a polynucleotide disclosed herein should provide a detection signal at least 5-, 10-, or 20-fold higher than the background hybridization provided with other unrelated sequences.

Detection of Expression Levels. Nucleotide probes are used to detect expression of a gene corresponding to the provided polynucleotide. In Northern blots, mRNA is separated electrophoretically and contacted with a probe. A probe is detected as hybridizing to an mRNA species of a particular size. The amount of hybridization is quantitated to determine relative amounts of expression, for example under a particular condition. Probes are used for in situ hybridization to cells to detect expression. Probes can also be used in vivo for diagnostic detection of hybridizing sequences.

Probes are typically labeled with a radioactive isotope. Other types of detectable labels can be used

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such as chromophores, fluors, and enzymes. Other examples of nucleotide hybridization assays are described in WO92/02526 and USPN 5,124,246.

Alternatively, the Polymerase Chain Reaction (PCR) is another means for detecting small amounts of target nucleic acids (see, e.g., Mullis et al., Meth. Enzymol. (1987) 155:335; USPN 4,683,195; and USPN 4,683,202). Two primer polymucleotides nucleotides that hybridize with the target nucleic acids are used to prime the reaction. The primers can be composed of sequence within or 3' and 5' to the polymucleotides of the Sequence Listing. Alternatively, if the primers are 3' and 5' to these polymucleotides, they need not hybridize to them or the complements. After amplification of the target with a thermostable polymerase, the amplified target nucleic acids can be detected by methods known in the art, e.g., Southern blot. mRNA or cDNA can also be detected by traditional blotting techniques (e.g., Southern blot, Northern blot, etc.) described in Sambrook et al., "Molecular Cloning: A Laboratory Manual" (New York, Cold Spring Harbor Laboratory, 1989) (e.g., without PCR amplification). In general, mRNA or cDNA generated from mRNA using a polymerase enzyme can be purified and separated using gel electrophorosis, and transferred to a solid support, such as nitrocellulose. The solid support is exposed to a labeled probe, washed to remove any unhybridized probe, and duplexes containing the labeled probe are detected.

Mapping. Polynucleotides of the present invention can be used to identify a chromosome on which the corresponding gene resides. Such mapping can be useful in identifying the function of the polynucleotide-related gene by its proximity to other genes with known function. Function can also be assigned to the polynucleotide-related gene when particular syndromes or diseases map to the same chromosome. For example, use of polynucleotide probes in identification and quantification of nucleic acid sequence aberrations is described in USPN 5,783,387. An exemplary mapping method is fluorescence in situ hybridization (FISH), which facilitates comparative genomic hybridization to allow total genome assessment of changes in relative copy number of DNA sequences (see, e.g., Valdes et al., Methods in Molecular Biology (1997) 68:1). Polynucleotides can also be mapped to particular chromosomes using, for example, radiation hybrids or chromosome-specific hybrid panels. See Leach et al., Advances in Genetics, (1995) 33:63-99; Walter et al., Nature Genetics (1994) 7:22; Walter and Goodfellow, Trends in Genetics (1992) 9:352. Panels for radiation hybrid mapping are available from Research Genetics, Inc., Huntsville, Alabama, USA. Databases for markers using various panels are available via the world wide web at sites supported by the Stanford Human Genome Center (Stanford University) and the Whitehead Institute for Biomedical Research/MIT Center for Genome Research. The statistical program RHMAP can be used to construct a map based on the data from radiation hybridization with a measure of the relative likelihood of one order versus another. RHMAP is available via the world wide web at a site supported by the University of Michigan. In addition,

commercial programs are available for identifying regions of chromosomes commonly associated with disease, such as cancer.

Tissue Typing or Profiling. Expression of specific mRNA corresponding to the provided polymucleotides can vary in different cell types and can be tissue-specific. This variation of mRNA levels in different cell types can be exploited with nucleic acid probe assays to determine tissue types. For example, PCR, branched DNA probe assays, or blotting techniques utilizing nucleic acid probes substantially identical or complementary to polynucleotides listed in the Sequence Listing can determine the presence or absence of the corresponding cDNA or mRNA.

Tissue typing can be used to identify the developmental organ or tissue source of a metastatic lesion by identifying the expression of a particular marker of that organ or tissue. If a polymeleotide is expressed only in a specific tissue type, and a metastatic lesion is found to express that polymeleotide, then the developmental source of the lesion has been identified. Expression of a particular polymeleotide can be assayed by detection of either the corresponding mRNA or the protein product. As would be readily apparent to any forensic scientist, the sequences disclosed herein are useful in differentiating human tissue from non-human tissue. In particular, these sequences are useful to differentiate human tissue from bird reptile, and amphibian tissue, for example.

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<u>Use of Polymorphisms</u>. A polymucleotide of the invention can be used in forensics, genetic analysis, mapping, and diagnostic applications where the corresponding region of a gene is polymorphic in the human population. Any means for detecting a polymorphism in a gene can be used, including, but not limited to electrophoresis of protein polymorphic variants, differential sensitivity to restriction enzyme cleavage, and hybridization to allele-specific probes.

Antibody Production. Expression products of a polynucleotide of the invention, as well as the corresponding mRNA, cDNA, or complete gene, can be prepared and used for raising antibodies for experimental, diagnostic, and therapeutic purposes. For polynucleotides to which a corresponding gene has not been assigned, this provides an additional method of identifying the corresponding gene. The polynucleotide or related cDNA is expressed as described above, and antibodies are prepared. These antibodies are specific to an epitope on the polypeptide encoded by the polynucleotide, and can precipitate or bind to the corresponding native protein in a cell or tissue preparation or in a cell-free extract of an in vitro expression system.

Methods for production of antibodies that specifically bind a selected antigen are well known in the art. Immunogens for raising antibodies can be prepared by mixing a polypeptide encoded by a polymeleotide of the invention with an adjuvant, and/or by making fusion proteins with larger immunogenic proteins. Polypeptides can also be covalently linked to other larger immunogenic proteins, such as keyhole limpet hemocyanin. Immunogens are typically administered intradermally,

subcutaneously, or intramuscularly to experimental animals such as rabbits, sheep, and mice, to generate antibodies. Monoclonal antibodies can be generated by isolating spleen cells and fissing myeloma cells to form hybridomas. Alternatively, the selected polynucleotide is administered directly, such as by intramuscular injection, and expressed in vivo. The expressed protein generates a variety of protein-specific immune responses, including production of antibodies, comparable to administration of the protein.

Preparations of polyalonal and monoclonal antibodies specific for polyapeptides encoded by a selected polynucleotide are made using standard methods known in the art. The antibodies specifically bind to epitopes present in the polypeptides encoded by polynucleotides disclosed in the Sequence Listing. Typically, at least 6, 8, 10, or 12 contiguous amino acids are required to form an epitope. Epitopes that involve non-contiguous amino acids may require a longer polypeptide, e.g., at least 15, 25, or 50 amino acids. Antibodies that specifically bind to human polypeptides encoded by the provided polypeptides should provide a detection signal at least 5-, 10-, or 20-fold higher than a detection signal provided with other proteins when used in Western blots or other immunochemical assays. Preferably, antibodies that specifically bind polypeptides contemplated by the invention do not bind to other proteins in immunochemical assays at detectable levels and can immunoprecipitate the specific polypeptide from solution.

The invention also contemplates naturally occurring antibodies specific for a polypeptide of the invention. For example, serum antibodies to a polypeptide of the invention in a human population can be purified by methods well known in the art, e.g., by passing antiserum over a column to which the corresponding selected polypeptide or fusion protein is bound. The bound antibodies can then be eluted from the column, for example, using a buffer with a high salt concentration.

In addition to the antibodies discussed above, the invention also contemplates genetically engineered antibodies entibodies (e.g., chimeric antibodies, humanized antibodies, human antibodies produced by a transgenic animal (e.g., a transgenic mouse such as the XenomousTM), antibody derivatives (e.g., single chain antibodies, antibody fragments (e.g., Fab, etc.)), according to methods well known in the art.

Polynucleotides or Arrays for Diagnostics.

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Polynucleotide arrays provide a high throughput technique that can assay a large number of polynucleotides in a sample. This technology can be used as a diagnostic and as tool to test for differential expression expression, e.g., to determine function of an encoded protein. A variety of methods of producing arrays, as well as variations of these methods, are known in the art and contemplated for use in the invention. For example, arrays can be created by spotting polynucleotide probes onto a substrate (e.g., slass, nitrocellulose, etc.) in a two-dimensional matrix or array having

bound probes. The probes can be bound to the substrate by either covalent bonds or by non-specific interactions, such as hydrophobic interactions. Samples of polymucleotides can be detectably labeled (e.g., using radioactive or fluorescent labels) and then hybridized to the probes. Double stranded polymucleotides, comprising the labeled sample polymucleotides bound to probe polymucleotides, can be detected once the unbound portion of the sample is washed away. Alternatively, the polymucleotides of the test sample can be immobilized on the array, and the probes detectably labeled. Techniques for constructing arrays and methods of using these arrays are described in, for example, Schena et al. (1996) Proc Natl Acad Sci U S A. 93(20):10614-9; Schena et al. (1995) Science 270(5235):467-70; Shalon et al. (1996) Genome Res. 6(7):639-45, USPN 5,807,522, EP 799 897; WO 97/22212; WO 97/227317; EP 785 280; WO 97/22357; USPN 5,593,839; USPN 5,578,832; EP 728 520; USPN 5,599,695; EP 721 016; USPN 5,556,752; WO 95/22058; and USPN 5,631,734.

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Arrays can be used to, for example, examine differential expression of genes and can be used to determine gene function. For example, arrays can be used to detect differential expression of a gene corresponding to a polymolectide of the invention, where expression is compared between a test cell and control cell (e.g., cancer cells and normal cells). For example, high expression of a particular message in a cancer cell, which is not observed in a corresponding normal cell, can indicate a cancer specific gene product. Exemplary uses of arrays are further described in, for example, Pappalarado et al., Sem. Radiation Oncol. (1998) 8:217; and Ramsay Nature Biotechnol. (1998) 16:40. Furthermore, many variations on methods of detection using arrays are well within the skill in the art and within the scope of the present invention. For example, rather than immobilizing the probe to a solid support, the test sample can be immobilized on a solid support which is then contacted with the probe.

Differential Expression in Diagnosis

The polynucleotides of the invention can also be used to detect differences in expression levels between two cells, e.g., as a method to identify abnormal or diseased tissue in a human. For polynucleotides corresponding to profiles of protein families, the choice of tissue can be selected according to the putative biological function. In general, the expression of a gene corresponding to a specific polynucleotide is compared between a first tissue that is suspected of being diseased and a second, normal tissue of the human. The tissue suspected of being abnormal or diseased can be derived from a different tissue type of the human, but preferably it is derived from the same tissue type; for example, an intestinal polyp or other abnormal growth should be compared with normal intestinal tissue. The normal tissue can be the same tissue as that of the test sample, or any normal tissue of the patient, especially those that express the polynucleotide-related gene of interest (e.g., brain, thymus, testis, heart, prostate, placenta, spleen, small intestine, skeletal muscle, pancreas, and the mucosal lining of the colon). A difference between the polynucleotide-related gene, mRNA, or protein in the

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two tissues which are compared, for example, in molecular weight, amino acid or nucleotide sequence, or relative abundance, indicates a change in the gene, or a gene which regulates it, in the tissue of the human that was suspected of being diseased. Examples of detection of differential expression and its use in diagnosis of cancer are described in USPNs 5,688,641 and 5,677,125.

A genetic predisposition to disease in a human can also be detected by comparing expression levels of an mRNA or protein corresponding to a polynucleotide of the invention in a fetal tissue with levels associated in normal fetal tissue. Fetal tissues that are used for this purpose include, but are not limited to, amniotic fluid, chorionic villi, blood, and the blastomere of an in vitro-fertilized embryo. The comparable normal polynucleotide-related gene is obtained from any tissue. The mRNA or protein is obtained from a normal tissue of a human in which the polynucleotide-related gene is expressed. Differences such as alterations in the nucleotide sequence or size of the same product of the fetal polynucleotide-related gene or mRNA, or alterations in the molecular weight, amino acid sequence, or relative abundance of fetal protein, can indicate a germline mutation in the polynucleotiderelated gene of the fetus, which indicates a genetic predisposition to disease. In general, diagnostic, prognostic, and other methods of the invention based on differential expression involve detection of a level or amount of a gene product, particularly a differentially expressed gene product, in a test sample obtained from a patient suspected of having or being susceptible to a disease (e.g., breast cancer, lung cancer, colon cancer and/or metastatic forms thereof), and comparing the detected levels to those levels found in normal cells (e.g., cells substantially unaffected by cancer) and/or other control cells (e.g., to differentiate a cancerous cell from a cell affected by dysplasia). Furthermore, the severity of the disease can be assessed by comparing the detected levels of a differentially expressed gene product with those levels detected in samples representing the levels of differentially expressed gene product associated with varying degrees of severity of disease. It should be noted that use of the term "diagnostic" herein is not necessarily meant to exclude "prognostic" or "prognosis," but rather is used as a matter of convenience.

The term "differentially expressed gene" is generally intended to encompass a polynucleotide that can, for example, include an open reading frame encoding a gene product (e.g., a polypeptide), and/or introns of such genes and adjacent 5' and 3' non-coding nucleotide sequences involved in the regulation of expression, up to about 20 kb beyond the coding region, but possibly further in either direction. The gene can be introduced into an appropriate vector for extrachromosomal maintenance or for integration into a host genome. In general, a difference in expression level associated with a decrease in expression level of at least about 25%, usually at least about 50% to 75%, more usually at least about 90% or more is indicative of a differentially expressed gene of interest, i.e., a gene that is underexpressed or down-regulated in the test sample relative to a control sample. Furthermore, a

difference in expression level associated with an increase in expression of at least about 25%, usually at least about 50% to 75%, more usually at least about 90% and can be at least about 1½-fold, usually at least about 2-fold to about 10-fold, and can be about 100-fold to about 1,000-fold increase relative to a control sample is indicative of a differentially expressed gene of interest, i.e., an overexpressed or up-regulated gene.

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"Differentially expressed polymucleotide" as used herein means a nucleic acid molecule (RNA or DNA) comprising a sequence that represents a differentially expressed gene, e.g., the differentially expressed polymucleotide comprises a sequence (e.g., an open reading frame encoding a gene product) that uniquely identifies a differentially expressed gene so that detection of the differentially expressed polymucleotide in a sample is correlated with the presence of a differentially expressed gene in a sample. "Differentially expressed polymucleotide" is also meant to encompass fragments of the disclosed polymucleotides, e.g., fragments retaining biological activity, as well as nucleic acids homologous, substantially similar, or substantially identical (e.g., having about 90% sequence identity) to the disclosed polynucleotides.

Methods of the subject invention useful in diagnosis or prognosis typically involve comparison of the abundance of a selected differentially expressed gene product in a sample of interest with that of a control to determine any relative differences in the expression of the gene product, where the difference can be measured qualitatively and/or quantitatively. Quantitation can be accomplished, for example, by comparing the level of expression product detected in the sample with the amounts of product present in a standard curve. A comparison can be made visually; by using a technique such as densitometry, with or without computerized assistance; by preparing a representative library of cDNA clones of mRNA isolated from a test sample, sequencing the clones in the library to determine that number of cDNA clones corresponding to the same gene product, and analyzing the number of clones corresponding to that same gene product relative to the number of clones of the same gene product in a control sample; or by using an array to detect relative levels of hybridization to a selected sequence or set of sequences, and comparing the hybridization pattern to that of a control. The differences in expression are then correlated with the presence or absence of an abnormal expression pattern. A variety of different methods for determining the nucleic acid abundance in a sample are known to those of skill in the art (see, e.g., WO 97/27317).

In general, diagnostic assays of the invention involve detection of a gene product of a polynucleotide sequence (e.g., mRNA or polypeptide) that corresponds to a sequence of SEQ ID NOS:1-6010. The patient from whom the sample is obtained can be apparently healthy, susceptible to disease (e.g., as determined by family history or exposure to certain environmental factors), or can

already be identified as having a condition in which altered expression of a gene product of the invention is implicated.

Diagnosis can be determined based on detected gene product expression levels of a gene product encoded by at least one, preferably at least two or more, at least 3 or more, or at least 4 or more of the polynucleotides having a sequence set forth in SEQ ID NOS:1-6010, and can involve detection of expression of genes corresponding to all of SEQ ID NOS:1-6010 and can involve detection of expression of genes corresponding to all of SEQ ID NOS:1-6010 additional sequences that can serve as additional diagnostic markers and/or reference sequences. Where the diagnostic method is designed to detect the presence or susceptibility of a patient to cancer, the assay preferably involves detection of a gene product encoded by a gene corresponding to a polynucleotide that is differentially expressed in cancer. Examples of such differentially expressed polynucleotides are described in the Examples below. Given the provided polynucleotides and information regarding their relative expression levels provided herein, assays using such polynucleotides and detection of their expression levels in diagnosis and prognosis will be readily apparent to the ordinarily skilled artisan.

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Any of a variety of detectable labels can be used in connection with the various embodiments of the diagnostic methods of the invention. Suitable detectable labels include fluorochromes, (e.g. fluorescein isothiocyanate (FITC), rhodamine, Texas Red, phycocrythrin, allophycocyanin, 6-carboxyfluorescein (6-FAM), 2',7'-dimethoxy-4',5'-dichloro-6-carboxyfluorescein, 6-carboxy-X-rhodamine (ROX), 6-carboxy-2',4',7',4,7-hexachlorofluorescein (HEX), 5-carboxyfluorescein (5-FAM) or N,N,N',N'-tetramethyl-6-carboxyrhodamine (TAMRA)), radioactive labels, (e.g. 32P, 35S, 3H, etc.), and the like. The detectable label can involve a two stage systems (e.g., biotin-avidin, hapten-anti-hapten anti-bapten anti-b

Reagents specific for the polymolectides and polypeptides of the invention, such as antibodies and nucleotide probes, can be supplied in a kit for detecting the presence of an expression product in a biological sample. The kit can also contain buffers or labeling components, as well as instructions for using the reagents to detect and quantify expression products in the biological sample. Exemplary embodiments of the diagnostic methods of the invention are described below in more detail.

Polypeptide detection in diagnosis. In one embodiment, the test sample is assayed for the level of a differentially expressed polypeptide. Diagnosis can be accomplished using any of a number of methods to determine the absence or presence or altered amounts of the differentially expressed polypeptide in the test sample. For example, detection can utilize staining of cells or histological sections with labeled antibodies, performed in accordance with conventional methods. Cells can be permeabilized to stain cytoplasmic molecules. In general, antibodies that specifically bind a differentially expressed polypeptide of the invention are added to a sample, and incubated for a period of time sufficient to allow binding to the epitope, usually at least about 10 minutes. The antibody can

be detectably labeled for direct detection (e.g., using radioisotopes, enzymes, fluorescers, chemiluminescers, and the like), or can be used in conjunction with a second stage antibody or reagent to detect binding (e.g., biotin with horseradish peroxidase-conjugated avidin, a secondary antibody conjugated to a fluorescent compound, e.g. fluorescein, rhodamine, Texas red, etc.). The absence or presence of antibody binding can be determined by various methods, including flow cytometry of dissociated cells, microscopy, radiography, scintillation counting, etc. Any suitable alternative methods of qualitative or quantitative detection of levels or amounts of differentially expressed polypeptide can be used, for example, ELISA, western blot, immunoprecipitation, radioimmunoassay, etc.

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mRNA detection. The diagnostic methods of the invention can also or alternatively involve detection of mRNA encoded by a gene corresponding to a differentially expressed polymeleotide of the invention. Any suitable qualitative or quantitative methods known in the art for detecting specific mRNAs can be used. mRNA can be detected by, for example, in situ hybridization in tissue sections, by reverse transcriptase-PCR, or in Northern blots containing poly A+mRNA. One of skill in the art can readily use these methods to determine differences in the size or amount of mRNA transcripts between two samples. mRNA expression levels in a sample can also be determined by generation of a library of expressed sequence tags (ESTs) from the sample, where the EST library is representative of sequences present in the sample (Adams, et al., (1991) Science 252:1651). Enumeration of the relative representation of ESTs within the library can be used to approximate the relative representation of the gene transcript within the starting sample. The results of EST analysis of a test sample can then be compared to EST analysis of a reference sample to determine the relative expression levels of a selected polynucleotide, particularly a polynucleotide corresponding to one or more of the differentially expressed genes described herein. Alternatively, gene expression in a test sample can be performed using serial analysis of gene expression (SAGE) methodology (e.g., Velculescu et al., Science (1995) 270:484) or differential display (DD) methodology (see, e.g., USPN 5,776.683 and USPN 5,807.680).

Alternatively, gene expression can be analyzed using hybridization analysis. Oligonucleotides or cDNA can be used to selectively identify or capture DNA or RNA of specific sequence composition, and the amount of RNA or cDNA hybridized to a known capture sequence determined qualitatively or quantitatively, to provide information about the relative representation of a particular message within the pool of cellular messages in a sample. Hybridization analysis can be designed to allow for concurrent screening of the relative expression of hundreds to thousands of genes by using, for example, array-based technologies having high density formats, including filters, microscope slides, or microchips, or solution-based technologies that use spectroscopic analysis (e.g., mass spectrometry).

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One exemplary use of arrays in the diagnostic methods of the invention is described below in more detail.

Use of a single gene in diagnostic applications. The diagnostic methods of the invention can focus on the expression of a single differentially expressed gene. For example, the diagnostic method can involve detecting a differentially expressed gene, or a polymorphism of such a gene (e.g., a polymorphism in a coding region or control region), that is associated with disease. Disease-associated polymorphisms can include deletion or truncation of the gene, mutations that alter expression level and/or affect activity of the encoded protein, etc.

A number of methods are available for analyzing nucleic acids for the presence of a specific sequence, e.g. a disease associated polymorphism. Where large amounts of DNA are available, genomic DNA is used directly. Alternatively, the region of interest is cloned into a suitable vector and grown in sufficient quantity for analysis. Cells that express a differentially expressed gene can be used as a source of mRNA, which can be assayed directly or reverse transcribed into cDNA for analysis. The nucleic acid can be amplified by conventional techniques, such as the polymerase chain reaction (PCR), to provide sufficient amounts for analysis, and a detectable label can be included in the amplification reaction (e.g., using a detectably labeled primer or detectably labeled oligonucleotides) to facilitate detection. Alternatively, various methods are also known in the art that utilize oligonucleotide ligation as a means of detecting polymorphisms, see, e.g., Riley et al., Nucl. Acids Res. (1990) 18:2887; and Delahunty et al., Am. J. Hum. Genet. (1996) 58:1239.

The amplified or cloned sample nucleic acid can be analyzed by one of a number of methods known in the art. The nucleic acid can be sequenced by dideoxy or other methods, and the sequence of bases compared to a selected sequence, e.g., to a wild-type sequence. Hybridization with the polymorphic or variant sequence can also be used to determine its presence in a sample (e.g., by Southern blot, dot blot, etc.). The hybridization pattern of a polymorphic or variant sequence and a control sequence to an array of oligonucleotide probes immobilized on a solid support, as described in US 5,445,934, or in WO 95/35505, can also be used as a means of identifying polymorphic or variant sequences associated with disease. Single strand conformational polymorphism (SSCP) analysis, denaturing gradient gel electrophoresis (DGGE), and heteroduplex analysis in gel matrices are used to detect conformational changes created by DNA sequence variation as alterations in electrophoretic mobility. Alternatively, where a polymorphism creates or destroys a recognition site for a restriction endonuclease, the sample is digested with that endonuclease, and the products size fractionated to determine whether the fragment was digested. Fractionation is performed by gel or capillary electrophoresis, particularly acrylamide or agarose gels.

Screening for mutations in a gene can be based on the functional or antigenic characteristics of the protein. Protein truncation assays are useful in detecting deletions that can affect the biological activity of the protein. Various immunoassays designed to detect polymorphisms in proteins can be used in screening. Where many diverse genetic mutations lead to a particular disease phenotype, functional protein assays have proven to be effective screening tools. The activity of the encoded protein can be determined by comparison with the wild-type protein.

Diagnosis, Prognosis, Assessment of Therapy (Therametrics), and Management of Cancer

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The polynucleotides of the invention, as well as their gene products, are of particular interest as genetic or biochemical markers (e.g., in blood or tissues) that will detect the earliest changes along the carcinogenesis pathway and/or to monitor the efficacy of various therapies and preventive interventions. For example, the level of expression of certain polynucleotides can be indicative of a poorer prognosis, and therefore warrant more aggressive chemo- or radio-therapy for a patient or vice versa. The correlation of novel surrogate tumor specific features with response to treatment and outcome in patients can define prognostic indicators that allow the design of tailored therapy based on the molecular profile of the tumor. These therapies include antibody targeting, antagonists (e.g., small molecules), and gene therapy. Determining expression of certain polynucleotides and comparison of a patient's profile with known expression in normal tissue and variants of the disease allows a determination of the best possible treatment for a patient, both in terms of specificity of treatment and in terms of comfort level of the patient. Surrogate tumor markers, such as polynucleotide expression, can also be used to better classify, and thus diagnose and treat, different forms and disease states of cancer. Two classifications widely used in oncology that can benefit from identification of the expression levels of the genes corresponding to the polynucleotides of the invention are staging of the cancerous disorder, and grading the nature of the cancerous tissue.

The polynucleotides that correspond to differentially expressed genes, as well as their encoded gene products, can be useful to monitor patients having or susceptible to cancer to detect potentially malignant events at a molecular level before they are detectable at a gross morphological level. In addition, the polynucleotides of the invention, as well as the genes corresponding to such polynucleotides, can be useful as therametrics, e.g., to assess the effectiveness of therapy by using the polynucleotides or their encoded gene products, to assess, for example, tumor burden in the patient before, during, and after therapy.

Furthermore, a polynucleotide identified as corresponding to a gene that is differentially expressed in, and thus is important for, one type of cancer can also have implications for development or risk of development of other types of cancer, e.g., where a polynucleotide represents a gene differentially expressed across various cancer types. Thus, for example, expression of a polynucleotide

corresponding to a gene that has clinical implications for metastatic colon cancer can also have clinical implications for stomach cancer or endometrial cancer.

Staging. Staging is a process used by physicians to describe how advanced the cancerous state is in a patient. Staging assists the physician in determining a prognosis, planning treatment and evaluating the results of such treatment. Staging systems vary with the types of cancer, but generally involve the following "TNM" system: the type of tumor, indicated by T; whether the cancer has metastasized to nearby lymph nodes, indicated by N; and whether the cancer has metastasized to more distant parts of the body, indicated by M. Generally, if a cancer is only detectable in the area of the primary lesion without having spread to any lymph nodes it is called Stage I. If it has spread only to the closest lymph nodes, it is called Stage II. In Stage III, the cancer has generally spread to the lymph nodes in near proximity to the site of the primary lesion. Cancers that have spread to a distant part of the body, such as the liver, bone, brain or other site, are Stage IV, the most advanced stage.

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The polynucleotides of the invention can facilitate fine-tuning of the staging process by identifying markers for the aggresivity of a cancer, o.g., the metastatic potential, as well as the presence in different areas of the body. Thus, a Stage II cancer with a polynucleotide signifying a high metastatic potential cancer can be used to change a borderline Stage II tumor to a Stage III tumor, justifying more aggressive therapy. Conversely, the presence of a polynucleotide signifying a lower metastatic potential allows more conservative staging of a tumor.

Grading of cancers. Grade is a term used to describe how closely a tumor resembles normal tissue of its same type. The microscopic appearance of a tumor is used to identify tumor grade based on parameters such as cell morphology, cellular organization, and other markers of differentiation. As a general rule, the grade of a tumor corresponds to its rate of growth or aggressiveness, with undifferentiated or high-grade tumors being more aggressive than well-differentiated or low-grade tumors. The following guidelines are generally used for grading tumors: 1) GX Grade cannot be assessed; 2) G1 Well differentiated; 3) G2 Moderately well differentiated; 4) G3 Poorly differentiated; 5) G4 Undifferentiated. The polymucleotides of the invention can be especially valuable in determining the grade of the tumor, as they not only can aid in determining the differentiation status of the cells of a tumor, they can also identify factors other than differentiation that are valuable in determining the aggressiveness of a tumor, such as metastatic potential.

<u>Detection of colon cancer</u>. The polymocleotides corresponding to genes that exhibit the appropriate expression pattern can be used to detect colon cancer in a subject. Colorectal cancer is one of the most common neoplasms in humans and perhaps the most frequent form of hereditary neoplasia. Prevention and early detection are key factors in controlling and curing colorectal cancer. Colorectal cancer begins as polyps, which are small, benign growths of cells that form on the inner lining of the

colon. Over a period of several years, some of these polyps accumulate additional mutations and become cancerous. Multiple familial colorectal cancer disorders have been identified, which are summarized as follows: 1) Familial adenomatous polyposis (FAP); 2) Gardner's syndrome; 3) Hereditary nonpolyposis colon cancer (HNPCC); and 4) Familial colorectal cancer in Ashkenazi Jews. The expression of appropriate polynucleotides of the invention can be used in the diagnosis, prognosis and management of colorectal cancer. Detection of colon cancer can be determined using expression levels of any of these sequences alone or in combination with the levels of expression. Determination of the aggressive nature and/or the metastatic potential of a colon cancer can be determined by comparing levels of one or more polynucleotides of the invention and comparing total levels of another sequence known to vary in cancerous tissue, e.g., expression of p53, DCC ras, lor FAP (see, e.g., Fearon ER, et al., Cell (1990) 61(5):759; Hamilton SR et al., Cancer (1993) 72:957; Bodmer W, et al., Nat Genet. (1994) 4(3):217; Fearon ER, Ann N Y Acad Sci. (1995) 768:101). For example, development of colon cancer can be detected by examining the ratio of any of the polynucleotides of the invention to the levels of oncogenes (e.g., ras) or tumor suppressor genes (e.g., FAP or p53). Thus, expression of specific marker polynucleotides can be used to discriminate between normal and cancerous colon tissue, to discriminate between colon cancers with different cells of origin, to discriminate between colon cancers with different potential metastatic rates, etc. For a review of markers of cancer, see, e.g., Hanahan et al. (2000) Cell 100:57-70.

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<u>Detection of prostate cancer.</u> The polynucleotides and their corresponding genes and gene products exhibiting the appropriate differential expression pattern can be used to detect prostate cancer in a subject. Over 95% of primary prostate cancers are adenocarcinomas. Signs and symptoms may include: frequent urination, especially at night; inability to urinate; trouble starting or holding back urination; a weak or interrupted urine flow; and frequent pain or stiffness in the lower back, hips or upper thiels.

Many of the signs and symptoms of prostate cancer can be caused by a variety of other noncancerous conditions. For example, one common cause of many of these signs and symptoms is a condition called benign prostatic hypertrophy, or BPH. In BPH, the prostate gets bigger and may block the flow of urine or interfere with sexual function. The methods and compositions of the invention can be used to distinguish between prostate cancer and such non-cancerous conditions. The methods of the invention can be used in conjunction with conventional methods of diagnosis, e.g., digital rectal exam and/or detection of the level of prostate specific antigen (PSA), a substance produced and secreted by the prostate.

<u>Detection of breast cancer</u>, The majority of breast cancers are adenocarcinoma subtypes, which can be summarized as follows: 1) ductal carcinoma in situ (DCIS), including comedocarcinoma;

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2) infiltrating (or invasive) ductal carcinoma (IDC); 3) lobular carcinoma in situ (LCIS); 4) infiltrating (or invasive) lobular carcinoma (ILC); 5) inflammatory breast cancer; 6) medullary carcinoma; 7) mucinous carcinoma; 8) Paget's disease of the nipple; 9) Phyllodes tumor; and 10) tubular carcinoma:

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The expression of polynucleotides of the invention can be used in the diagnosis and management of breast cancer, as well as to distinguish between types of breast cancer. Detection of breast cancer can be determined using expression levels of any of the appropriate polynucleotides of the invention, either alone or in combination. Determination of the aggressive nature and/or the metastatic potential of a breast cancer can also be determined by comparing levels of one or more polynucleotides of the invention and comparing levels of another sequence known to vary in cancerous tissue, e.g., ER expression. In addition, development of breast cancer can be detected by examining the ratio of expression of a differentially expressed polynucleotide to the levels of steroid hormones (e.g., testosterone or estrogen) or to other hormones (e.g., growth hormone, insulin). Thus, expression of specific marker polynucleotides can be used to discriminate between normal and cancerous breast tissue, to discriminate between breast cancers with different cells of origin, to discriminate between breast cancers with different potential metastatic rates, etc.

Detection of lung cancer. The polynucleotides of the invention can be used to detect lung cancer in a subject. Although there are more than a dozen different kinds of lung cancer, the two main types of lung cancer are small cell and nonsmall cell, which encompass about 90% of all lung cancer cases. Small cell carcinoma (also called oat cell carcinoma) usually starts in one of the larger bronchial tubes, grows fairly rapidly, and is likely to be large by the time of diagnosis. Nonsmall cell lung cancer (NSCLC) is made up of three general subtypes of lung cancer. Epidermoid carcinoma (also called squamous cell carcinoma) usually starts in one of the larger bronchial tubes and grows relatively slowly. The size of these tumors can range from very small to quite large. Adenocarcinoma starts growing near the outside surface of the lung and can vary in both size and growth rate. Some slowly growing adenocarcinomas are described as alveolar cell cancer. Large cell carcinoma starts near the surface of the lung, grows rapidly, and the growth is usually fairly large when diagnosed. Other less common forms of lung cancer are carcinoid, cylindroma, mucoepidermoid, and malignant mesothelioma.

The polynucleotides of the invention, e.g., polynucleotides differentially expressed in normal cells versus cancerous lung cells (e.g., tumor cells of high or low metastatic potential) or between types of cancerous lung cells (e.g., high metastatic versus low metastatic), can be used to distinguish types of lung cancer as well as identifying traits specific to a certain patient's cancer and selecting an appropriate therapy. For example, if the patient's biopsy expresses a polynucleotide that is associated with a low metastatic potential, it may justify leaving a larger portion of the patient's lung in surgery to

remove the lesion. Alternatively, a smaller lesion with expression of a polynucleotide that is associated with high metastatic potential may justify a more radical removal of lung tissue and/or the surrounding lymph nodes, even if no metastasis can be identified through pathological examination.

Use of Polynucleotides to Screen for Peptide Analoes and Antaeonists

Polypeptides encoded by the instant polynucleotides and corresponding full-length genes can be used to screen peptide libraries to identify binding partners, such as receptors, from among the encoded polypeptides. Peptide libraries can be synthesized according to methods known in the art (see, e.g., USPN 5,010,175, and WO 91/17823). Agonists or antagonists of the polypeptides of the invention can be screened using any available method known in the art, such as signal transduction, antibody binding, receptor binding, mitogenic assays, chemotaxis assays, etc. The assay conditions ideally should resemble the conditions under which the native activity is exhibited in vivo, that is, under physiologic pH, temperature, and ionic strength. Suitable agonists or antagonists will exhibit strong inhibition or enhancement of the native activity at concentrations that do not cause toxic side effects in the subject. Agonists or antagonists that compete for binding to the native polypeptide can require concentrations equal to or greater than the native concentration, while inhibitors capable of binding irreversibly to the polypeptide can be added in concentrations on the order of the native concentration.

Such screening and experimentation can lead to identification of a novel polypeptide binding partner, such as a receptor, encoded by a gene or a cDNA corresponding to a polynucleotide of the invention, and at least one peptide agonist or antagonist of the novel binding partner. Such agonists and antagonists can be used to modulate, enhance, or inhibit receptor function in cells to which the receptor is native, or in cells that possess the receptor as a result of genetic engineering. Further, if the novel receptor shares biologically important characteristics with a known receptor, information about agonist/antagonist binding can facilitate development of improved agonists/antagonists of the known receptor.

25 Pharmaceutical Compositions and Uses

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Pharmaceutical compositions can comprise polypeptides, receptors that specifically bind a polypeptide produced by a differentially expressed gene (e.g., antibodies, or polynucleotides (including antisense nucleotides and ribozymes) of the claimed invention in a therapeutically effective amount. The compositions can be used to treat primary tumors as well as metastases of primary tumors. In addition, the pharmaceutical compositions can be used in conjunction with conventional methods of cancer treatment, e.g., to sensitize tumors to radiation or conventional chemotherapy.

Where the pharmaccutical composition comprises a receptor (such as an antibody) that specifically binds to a gene product encoded by a differentially expressed gene, the receptor can be coupled to a drug for delivery to a treatment site or coupled to a detectable label to facilitate imaging of

a site comprising colon cancer cells. Methods for coupling antibodies to drugs and detectable labels are well known in the art, as are methods for imaging using detectable labels.

The term "therapeutically effective amount" as used herein refers to an amount of a therapeutic agent to treat, ameliorate, or prevent a desired disease or condition, or to exhibit a detectable therapeutic or preventative effect. The effect can be detected by, for example, chemical markers or antigen levels. Therapeutic effects also include reduction in physical symptoms, such as decreased body temperature. The precise effective amount for a subject will depend upon the subject's size and health, the nature and extent of the condition, and the therapeutics or combination of therapeutics selected for administration. Thus, it is not useful to specify an exact effective amount in advance. However, the effective amount for a given situation is determined by routine experimentation and is within the judgment of the clinician. For purposes of the present invention, an effective dose will generally be from about 0.01 mg/kg to 50 mg/kg or 0.05 mg/kg to about 10 mg/kg of the DNA constructs in the individual to which it is administrated.

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Delivery Methods

A pharmaceutical composition can also contain a pharmaceutically acceptable carrier. The term "pharmaceutically acceptable carrier" refers to a carrier for administration of a therapeutic agent, such as antibodies or a polypeptide, genes, and other therapeutic agents. The term refers to any pharmaceutical carrier that does not itself induce the production of antibodies harmful to the individual receiving the composition, and which can be administered without undue toxicity. Suitable carriers can be large, slowly metabolized macromolecules such as proteins, polysaccharides, polylactic acids, polyglycolic acids, polymeric amino acids, amino acid copolymers, and inactive virus particles. Such carriers are well known to those of ordinary skill in the art. Pharmaceutically acceptable carriers in therapeutic compositions can include liquids such as water, saline, glycerol and ethanol. Auxiliary substances, such as wetting or emulsifying agents, pH buffering substances, and the like, can also be present in such vehicles. Typically, the therapeutic compositions are prepared as injectables, either as liquid solutions or suspensions; solid forms suitable for solution in, or suspension in, liquid vehicles prior to injection can also be prepared. Liposomes are included within the definition of a pharmaceutically acceptable carrier. Pharmaceutically acceptable salts can also be present in the pharmaceutical composition, e.g., mineral acid salts such as hydrochlorides, hydrobromides, phosphates, sulfates, and the like; and the salts of organic acids such as acetates, propionates, malonates, benzoates, and the like. A thorough discussion of pharmaceutically acceptable excipients is available in Remington's Pharmaceutical Sciences (Mack Pub. Co., N.J. 1991).

Once formulated, the compositions of the invention can be (1) administered directly to the subject (e.g., as polynucleotide or polypeptides); or (2) delivered ex vivo, to cells derived from the

subject (e.g., as in ex vivo gene therapy). Direct delivery of the compositions will generally be accomplished by parenteral injection, e.g., subcutaneously, intraperitoneally, intravenously or intramuscularly, intratumorally or to the interstitial space of a tissue. Other modes of administration include oral and pulmonary administration, suppositories, and transdermal applications, needles, and gene guns or hyposprays. Dosage treatment can be a single dose schedule or a multiple dose schedule.

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Methods for the ex vivo delivery and reimplantation of transformed cells into a subject are known in the art and described in, e.g., WO 93/14778. Examples of cells useful in ex vivo applications include, for example, stem cells, particularly hematopoetic, lymph cells, macrophages, dendritic cells, or tumor cells. Generally, delivery of nucleic acids for both ex vivo and in vitro applications can be accomplished by, for example, dextran-mediated transfection, calcium phosphate precipitation, polybrene mediated transfection, protoplast fusion, electroporation, encapsulation of the polynucleotide(s) in liposomes, and direct microinjection of the DNA into nuclei, all well known in the art.

Once differential expression of a gene corresponding to a polynucleotide of the invention has been found to correlate with a proliferative disorder, such as neoplasia, dysplasia, and hyperplasia, the disorder can be amenable to treatment by administration of a therapeutic agent based on the provided polynucleotide, corresponding polypeptide or other corresponding molecule (e.g., antisense, ribozyme, etc.). In other embodiments, the disorder can be amenable to treatment by administration of a small molecule drug that, for example, serves as an inhibitor (antagonist) of the function of the encoded gene product of a gene having increased expression in cancerous cells relative to normal cells or as an agonist for gene products that are decreased in expression in cancerous cells (e.g., to promote the activity of sene products that act as tumor suppressors).

The dose and the means of administration of the inventive pharmaceutical compositions are determined based on the specific qualities of the therapeutic composition, the condition, age, and weight of the patient, the progression of the disease, and other relevant factors. For example, administration of polynucleotide therapeutic composition agents of the invention includes local or systemic administration, including injection, oral administration, particle gun or catheterized administration, and topical administration. Preferably, the therapeutic polynucleotide composition contains an expression construct comprising a promoter operably linked to a polynucleotide of at least 12, 22, 25, 30, or 35 contiguous nt of the polynucleotide of the invention. Various methods can be used to administer the therapeutic composition directly to a specific site in the body. For example, a small metastatic lesion is located and the therapeutic composition injected several times in several different locations within the body of tumor. Alternatively, arteries that serve a tumor are identified, and the therapeutic composition injected into such an artery, in order to deliver the composition directly into

the tumor. A tumor that has a necrotic center is aspirated and the composition injected directly into the now empty center of the tumor. The antisense composition is directly administered to the surface of the tumor, for example, by topical application of the composition. X-ray imaging is used to assist in certain of the above delivery methods.

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Targeted delivery of therapeutic compositions containing an antisense polynucleotide, subgenomic polynucleotides, or antibodies to specific tissues can also be used. Receptor-mediated DNA delivery techniques are described in, for example, Findeis et al., Trends Biotechnol. (1993) 11:202; Chiou et al., Gene Therapeutics: Methods And Applications Of Direct Gene Transfer (J.A. Wolff. ed.) (1994); Wu et al., J. Biol. Chem. (1988) 263:621; Wu et al., J. Biol. Chem. (1994) 269:542; Zenke et al., Proc. Natl. Acad. Sci. (USA) (1990) 87:3655; Wu et al., J. Biol. Chem. (1991) 266:338. Therapeutic compositions containing a polynucleotide are administered in a range of about 100 ng to about 200 mg of DNA for local administration in a gene therapy protocol. Concentration ranges of about 500 ng to about 50 mg, about 1 micrograms to about 2 mg, about 5 micrograms to about 500 micrograms, and about 20 micrograms to about 100 micrograms of DNA can also be used during a gene therapy protocol. Factors such as method of action (e.g., for enhancing or inhibiting levels of the encoded gene product) and efficacy of transformation and expression are considerations which will affect the dosage required for ultimate efficacy of the antisense subgenomic polynucleotides. Where greater expression is desired over a larger area of tissue, larger amounts of antisense subgenomic polynucleotides or the same amounts readministered in a successive protocol of administrations, or several administrations to different adjacent or close tissue portions of, for example, a tumor site, may be required to effect a positive therapeutic outcome. In all cases, routine experimentation in clinical trials will determine specific ranges for optimal therapeutic effect. For polynucleotide related genes encoding polypeptides or proteins with anti-inflammatory activity, suitable use, doses, and administration are described in USPN 5.654.173.

The therapeutic polynucleotides and polypeptides of the present invention can be delivered using gene delivery vehicles. The gene delivery vehicle can be of viral or non-viral origin (see generally, Jolly, Cancer Gene Therapy (1994) 1:51; Kimura, Human Gene Therapy (1994) 5:845; Connelly, Human Gene Therapy (1995) 1:185; and Kaplitt, Nature Genetics (1994) 6:148). Expression of such coding sequences can be induced using endogenous mammalian or heterologous promoters. Expression of the coding sequence can be either constitutive or regulated.

Viral-based vectors for delivery of a desired polynucleotide and expression in a desired cell are well known in the art. Exemplary viral-based vehicles include, but are not limited to, recombinant retroviruses (see, e.g., WO 90/07936; WO 94/03622; WO 93/25698; WO 93/25234; USPN 5, 219.740; WO 93/11230; WO 93/10218; USPN 4.777.127; GB Patent No. 2.200.651; EP 0 345 242;

and WO 91/02805), alphavirus-based vectors (e.g., Sindbis virus vectors, Semliki forest virus (ATCC VR-67; ATCC VR-1247), Ross River virus (ATCC VR-373; ATCC VR-1246) and Venezuelan equine encephalitis virus (ATCC VR-923; ATCC VR-1250; ATCC VR 1249, ATCC VR-532), and adeno-associated virus (AAV) vectors (see, e.g., WO 94/12649, WO 93/03769; WO 93/19191; WO 94/28938; WO 95/11984 and WO 95/00655). Administration of DNA linked to killed adenovirus, as described in Curiel, Hum. Gene Ther. (1992) 3:147, can also be employed.

Non-viral delivery vehicles and methods can also be employed, including, but not limited to, polycationic condensed DNA linked or unlinked to killed adenovirus alone (see, e.g., Curiel, Hum. Gene Ther. (1992) 3:147); ligand-linked DNA (see, e.g., Wu, J. Biol. Chem. (1989) 264:16985); cukaryotic cell delivery vehicles cells (see, e.g., USPN 5,814,482; WO 95/07994; WO 96/17072; WO 95/030763; and WO 97/42338) and nucleic charge neutralization or fusion with cell membranes. Naked DNA can also be employed. Exemplary naked DNA introduction methods are described in WO 90/11092 and USPN 5,580,859. Liposomes that can act as gene delivery vehicles are described in USPN 5,422,120; WO 95/13796; WO 94/23697; WO 91/14445; and EP 0524968. Additional approaches are described in Philip, Mol. Cell Biol. (1994) 14:2411, and in Woffendin, Proc. Natl. Acad. Sci. (1994) 91:1581

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Further non-viral delivery suitable for use includes mechanical delivery systems such as the approach described in Woffendin et al., Proc. Natl. Acad. Sci. USA (1994) 91(24):11581. Moreover, the coding sequence and the product of expression of such can be delivered through deposition of photopolymerized hydrogel materials or use of ionizing radiation (see, e.g., USPN 5,206,152 and WO 92/11033). Other conventional methods for gene delivery that can be used for delivery of the coding sequence include, for example, use of hand-held gene transfer particle gun (see, e.g., USPN 5,149,655); use of ionizing radiation for activating transferred gene (see, e.g., USPN 5,206,152 and WO 92/11033).

The present invention will now be illustrated by reference to the following examples which set forth particularly advantageous embodiments. However, it should be noted that these embodiments are illustrative and are not to be construed as restricting the invention in any way.

EXAMPLES

The following examples are put forth so as to provide those of ordinary skill in the art with a complete disclosure and description of how to make and use the present invention, and are not intended to limit the scope of what the inventors regard as their invention nor are they intended to represent that the experiments below are all or the only experiments performed. It will be readily apparent to those skilled in the art that the formulations, dosages, methods of administration, and other parameters of

this invention may be further modified or substituted in various ways without departing from the spirit and scope of the invention. Efforts have been made to ensure accuracy with respect to numbers used (e.g. amounts, temperature, etc.) but some experimental errors and deviations should be accounted for. Unless indicated otherwise, parts are parts by weight, molecular weight is weight average molecular weight, temperature is in degrees Centigrade, and pressure is at or near atmospheric.

Example 1: Source of Biological Materials and Overview of Novel Polynucleotides Expressed by the Biological Materials

Candidate polynucleotides that may represent novel polynucleotides were obtained from cDNA libraries generated from selected cell lines and patient tissues. In order to obtain the candidate polynucleotides, mRNA was isolated from several selected cell lines and patient tissues, and used to construct cDNA libraries. The cells and tissues that served as sources for these cDNA libraries are summarized in Table 1 below.

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Human colon cancer cell line Km12L4-A (Morikawa, et al., Cancer Research (1988) 48;6863) is derived from the KM12C cell line, The KM12C cell line (Morikawa et al. Cancer Res. (1988) 48:1943-1948), which is poorly metastatic (low metastatic) was established in culture from a Dukes' stage B2 surgical specimen (Morikawa et al. Cancer Res. (1988) 48:6863). The KM12L4-A is a highly metastatic subline derived from KM12C (Yeatman et al. Nucl. Acids. Res. (1995) 23:4007; Bao-Ling et al, Proc. Annu. Meet. Am. Assoc. Cancer. Res. (1995) 21;3269). The KM12C and KM12C-derived cell lines (e.g., KM12L4, KM12L4-A, etc.) are well-recognized in the art as a model 20 cell line for the study of colon cancer (see, e.g., Moriakawa et al., supra; Radinsky et al. Clin. Cancer Res. (1995) 1:19; Yeatman et al., (1995) supra; Yeatman et al. Clin. Exp. Metastasis (1996) 14:246). The MDA-MB-231 cell line (Brinkley et al. Cancer Res. (1980) 40:3118-3129) was originally

isolated from pleural effusions (Cailleau, J. Natl. Cancer. Inst. (1974) 53:661), is of high metastatic potential, and forms poorly differentiated adenocarcinoma grade II in nude mice consistent with breast carcinoma. The MCF7 cell line was derived from a pleural effusion of a breast adenocarcinoma and is non-metastatic. The MV-522 cell line is derived from a human lung carcinoma and is of high metastatic potential. The UCP-3 cell line is a low metastatic human lung carcinoma cell line; the MV-522 is a high metastatic variant of UCP-3. These cell lines are well-recognized in the art as models for the study of human breast and lung cancer (see, e.g., Chandrasekaran et al., Cancer Res. (1979) 39:870 (MDA-MB-231 and MCF-7): Gastnar et al., J Med Chem (1998) 41:4965 (MDA-MB-231 and MCF-7); Ranson et al., Br J Cancer (1998) 77:1586 (MDA-MB-231 and MCF-7); Kuang et al., Nucleic Acids Res (1998) 26:1116 (MDA-MB-231 and MCF-7); Varki et al., Int J Cancer (1987) 40:46 (UCP-3); Varki et al., Tumour Biol. (1990) 11:327; (MV-522 and UCP-3); Varki et al., Anticancer

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Res. (1990) 10:637; (MV-522); Kelner et al., Anticancer Res (1995) 15:867 (MV-522); and Zhang et al., Anticancer Drugs (1997) 8:696 (MV522)).

The samples of libraries 15-20 are derived from two different patients (UC#2, and UC#3). The bFGF-treated HMVEC were prepared by incubation with bFGF at 10ng/ml for 2 hrs; the VEGF-treated HMVEC were prepared by incubation with VEGF for 2 hrs. Following incubation with the respective growth factor, the cells were washed and lysis buffer added for RNA preparation. The GRRpz and WOca cell lines were provided by Dr. Douna M. Pechl, Department of Medicine, Stanford University School of Medicine. GRRpz was derived from normal prostate epithelium. The WOca cell line is a Glesson Grade 4 cell line.

Table 1. Description of cDNA Libraries

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Library (lib#)	Description	Number o Clones in Library
0	Artificial library composed of deselected clones (clones with no associated variant or cluster)	673
1	Human Colon Cell Line Km12 L4: High Metastatic Potential (derived from Km12C)	308731
2	Human Colon Cell Line Km12C: Low Metastatic Potential	284771
3	Human Breast Cancer Cell Line MDA-MB-231: High Metastatic Potential; micro-mets in lung	326937
4	Human Breast Cancer Cell Line MCF7: Non Metastatic	318979
8	Human Lung Cancer Cell Line MV-522: High Metastatic Potential	223620
9	Human Lung Cancer Cell Line UCP-3: Low Metastatic Potential	312503
12	Human microvascular endothelial cells (HMEC) - UNTREATED (PCR (OligodT) cDNA library)	41938
13	Human microvascular endothelial cells (HMEC) - bFGF TREATED (PCR (OligodT) cDNA library)	42100
14	Human microvascular endothelial cells (HMEC) - VEGF TREATED (PCR (OligodT) cDNA library)	42825
15	Normal Colon - UC#2 Patient (MICRODISSECTED PCR (OligodT) cDNA library)	282722
16	Colon Tumor - UC#2 Patient (MICRODISSECTED PCR (OligodT) cDNA library)	298831
17	Liver Metastasis from Colon Tumor of UC#2 Patient (MICRODISSECTED PCR (OligodT) cDNA library)	303467
18	Normal Colon - UC#3 Patient (MICRODISSECTED PCR (OligodT) cDNA library)	36216
19	Colon Tumor - UC#3 Patient (MICRODISSECTED PCR (OligodT) cDNA library)	41388
20	Liver Metastasis from Colon Tumor of UC#3 Patient (MICRODISSECTED PCR (OligodT) cDNA library)	30956
21	GRRpz Cells derived from normal prostate epithelium	164801
22	WOca Cells derived from Gleason Grade 4 prostate cancer epithelium	162088
23	Normal Lung Epithelium of Patient #1006 (MICRODISSECTED PCR (OligodT) cDNA library)	306198

Library (lib#)	Description	Number of Clones in Library
24	Primary tumor, Large Cell Carcinoma of Patient #1006 (MICRODISSECTED PCR (OligodT) cDNA library)	309349
25	Normal Prostate Epithelium from Patient IF97-26811	279444
26	Prostate Cancer Epithelium Gleason 3+3 Patient IF97-26811	269406
27	Normal Breast Epithelium from Patient 515	239494
28	Primary Breast tumor from Patient 515	259960
29	Lymph node metastasis from Patient 515	326786
30	Normal Prostate Epithelium from Chiron Patient ID 884	298431
31	Prostate Cancer Epithelium (Gleason 4+4) from Chiron Patient ID 884	331941

Characterization of sequences in the libraries

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was ribosomal or vector-derived.

After using the software program Phred (ver 0.000925.c, Green and Weing, ©1993-2000) to select those polynucleotides having the best quality sequence, the polyneucleotides were compared against the public databases to identify any homolgous sequences. The sequences of the isolated polynucleotides were first masked to eliminate low complexity sequences using the BLASTX masking program (Claverie "Effective Large-Scale Sequence Similarity Searches," In: Computer Methods for Macromolecular Sequence Analysis, Doolittle, ed., Meth. Enzymol. 266:212-227 Academic Press, NY, NY (1996); see particularly Claverie, in "Automated DNA Sequencing and Analysis Techniques" Adams et al., eds., Chap. 36, p. 267 Academic Press, San Diego, 1994 and Claverie et al. Comput. Chem. (1993) 17:191). Generally, masking does not influence the final search results, except to eliminate sequences of relatively little interest due to their low complexity, and to eliminate multiple "hits" based on similarity to repetitive regions common to multiple sequences, e.g., Alu repeats. The remaining sequences were then used in a BLASTN vs. GenBank search; sequences that exhibited greater than 70% overlap, 99% identity, and a p value of less than 1 x 10e-40 were discarded. Sequences from this search also were discarded if the inclusive parameters were met, but the sequence

The resulting sequences from the previous search were classified into three groups (1, 2 and 3 below) and searched in a BLASTX vs. NRP (non-redundant proteins) database search: (1) unknown (no hits in the GenBank search), (2) weak similarity (greater than 45% identity and p value of less than 1 x 10e-5), and (3) high similarity (greater than 60% overlap, greater than 80% identity, and p value less than 1 x 10e-5). Sequences having greater than 70% overlap, greater than 99% identity, and p value of less than 1 x 10e-40 were discarded.

The remaining sequences were classified as unknown (no hits), weak similarity, and high

similarity (parameters as above). Two searches were performed on these sequences. First, a BLAST

vs. EST database search was performed and sequences with greater than 99% overlap, greater than

99% similarity and a p value of less than $1 \times 10e$ -40 were discarded. Sequences with a p value of less than $1 \times 10e$ -65 when compared to a database sequence of human origin were also excluded. Second, a BLASTN vs. Patent GeneSeq database was performed and sequences having greater than 99% identity, p value less than $1 \times 10e$ -40, and greater than 99% overlap were discarded.

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The remaining sequences were subjected to screening using other rules and redundancies in the dataset. Sequences with a p value of less than 1 x 10e-111 in relation to a database sequence of human origin were specifically excluded. The final result provided the 8064 sequences listed as SEQ ID NOS:1-6010 in the accompanying Sequence Listing and summarized in Table 2 (inserted prior to claims). Each identified polymelecotide represents sequence from at least a partial mRNA transcript. Summary of polymelecotides of the invention

Table 2 (inserted prior to claims) provides a summary of polymucleotides isolated as described. Specifically, Table 2 provides: 1) the SEQ ID NO ("SEQ ID") assigned to each sequence for use in the present specification; 2) the Cluster Identification No. ("CLUSTER"); 3) the Sequence Name assigned to each sequence; 3) the sequence name ("SEQ NAME") used as an internal identifier of the sequence; 4) the orientation of the sequence ("ORIENT") (either forward (F) or reverse (R)); 5) the name assigned to the clone from which the sequence was isolated ("CLONE ID"); and the name of the library from which the sequence was isolated ("CLONE ID"); and the name of the clone from which the sequence was isolated the sequence was isolated from normal colon tissue of the patient assigned the idnetification UC#2). Because at least some of the provided polymucleotides represent partial mRNA transcripts, two or more polymucleotides may represent different regions of the same mRNA transcript and the same gene and/or may be contained within the same clone. Thus, for example, if two or more SEQ ID NOS: are identified as belonging to the same clone, then either sequence can be used to obtain the full-leneth mRNA or sene

25 Example 2:Results of Public Database Search to Identify Function of Gene Products

SEQ ID NOS:1-6010 were translated in all three reading frames, and the nucleotide sequences and translated amino acid sequences used as query sequences to search for homologous sequences in either the GenBank (nucleotide sequences) or Non-Redundant Protein (amino acid sequences) databases. Query and individual sequences were aligned using the BLAST 2.0 programs, available over the world wide at a site sponsored by the National Center for Biotechnology Information, which is supported by the National Library of Medicine and the National Institutes of Health (see also Altschul, et al. Nucleic Acids Res. (1997) 25:3389-3402). The sequences were masked to various extents to prevent searching of repetitive sequences or poly-A sequences, using the BLASTX program for masking low complexity as described above in Example 1.

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Tables 3A and 3B (inserted prior to claims) provides the alignment summaries having a p value of 1 x 10e-2 or less indicating substantial homology between the sequences of the present invention and those of the indicated public databases. Specifically, Table 3A provides the SEO ID NO of the query sequence, the accession number of the GenBank database entry of the homologous sequence, and the individual p value of each alignment. Table 3A provides the SEQ ID NO of the query sequence, the accession number of the Non-Redundant Protein database entry of the homologous sequence, and the individual p value of each alignment. The alignments provided in Tables 3A and 3B are the best available alignment to a DNA or amino acid sequence at a time just prior to filing of the present specification. The activity of the polypeptide encoded by the SEO ID NOS listed in these tables can be extrapolated to be substantially the same or substantially similar to the activity of the reported nearest neighbor or closely related sequence. The accession number of the nearest neighbor is reported, providing a publicly available reference to the activities and functions exhibited by the nearest neighbor. The public information regarding the activities and functions of each of the nearest neighbor sequences is incorporated by reference in this application. Also incorporated by reference is all publicly available information regarding the sequence, as well as the putative and actual activities and functions of the nearest neighbor sequences listed in Tables 3A and 3B and their related sequences. The search program and database used for the alignment, as well as the calculation of the p value are also indicated.

Full length sequences or fragments of the polynucleotide sequences of the nearest neighbors can be used as probes and primers to identify and isolate the full length sequence of the corresponding polynucleotide. The nearest neighbors can indicate a tissue or cell type to be used to construct a library for the full-length sequences of the corresponding polynucleotides.

Example 3: Members of Protein Families

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SEQ ID NOS:1-6010 were used to conduct a profile search as described in the specification above. Several of the polynucleotides of the invention were found to encode polypeptides having characteristics of a polypeptide belonging to a known protein family (and thus represent members of these protein families) and/or comprising a known functional domain. Table 4 (inserted before claims) provides the SEQ ID NO: of the query sequence, the Sequence Name, the Cluster to which the sequence is assigned, a brief description of the profile hit, the orientation (Direction, "Dir") of the query sequence with respect to the individual sequence)where forward (for) indicates that the alignment is in the same direction (left to right) as the sequence provided in the Sequence Listing and reverse (rev) indicates that the alignment is with a sequence complementary to the sequence provided in the Sequence Listing), and the secre of the profile hit.

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Some polynucleotides exhibited multiple profile hits where the query sequence contains overlapping profile regions, and/or where the sequence contains two different functional domains. Each of the profile hits of Table 4 is described in more detail below. The acronyms for the profiles (provided in parentheses) are those used to identify the profile in the Pfam, Prosite, and InterPro databases. The Pfam database can be accessed through web sites supported by Genome Sequencing Center at the Washington University School of Medicine or by the European Molecular Biology Laboratories in Heidelberg, Germany. The Prosite database can be accessed at the ExPASy Molecular Biology Server on the internet. The InterPro database can be accessed at web site supported by the EMBL European Bioinformatics Institute. The public information available on the Pfam, Prosite, and InterPro databases regarding the various profiles, including but not limited to the activities, function, and consensus sequences of various proteins families and protein domains, is incorporated herein by reference.

Seven Transmembrane Integral Membrane Proteins -- Rhodopsin Family (7tm_1: Pfam Accession No. PF00001). SEQ ID NOS:2973, 5467, and 5508 correspond to sequences encoding a polypeptides that are members of the seven transmembrane (7tm) receptor rhodopsin family. G-protein coupled receptors of the (7tm) rhodopsin family (also called R7G) are an extensive group of hormones, neurotransmitters, and light receptors which transduce extracellular signals by interaction with guanine nucleotide-binding (G) proteins (Strosberg, Eur. J. Biochem. (1991) 196:1; Kerlavage, Curr. Opin. Struct. Biol. (1991) 1:394; Probst et al., DNA Cell Biol. (1992) 11:1; Savarese et al., Biochem. J. (1992) 283:1. The consensus pattern that contains the conserved triplet and that also spans the major part of the third transmembrane helix is used to detect this widespread family of proteins: [GSTALIVMFYWC]-[GSTANCPDE]-{EDPKRH}-x(2)-[LIVMNQGA]-x(2)-[LIVMFT]-[GSTANC]-[LIVMFYWSTAC]-[DENH]-R-FFYWCSH]-x(2)-[LIVMNQGA]-x(2)-[LIVMFT]-

Ank Repeats (ANK; Pfam Accession No. PF0023). SEQ ID NOS:445, 487 and 3013 represent polynucleotides encoding Ank repeat-containing proteins. The ankyrin motif is a 33 amino acid sequence named after the protein ankyrin which has 24 tandem 33-amino-acid motifs. Ank repeats were originally identified in the cell-cycle-control protein cdc10 (Breeden et al., Nature (1987) 329:651). Proteins containing ankyrin repeats include ankyrin, myotropin, I-kappaB proteins, cell cycle protein cdc10, the Notch receptor (Matsune et al., Development (1997) 124(21):4265); G9a (or BAT8) of the class III region of the major histocompatibility complex (Biochem J. (1993) 290:811-818); FABP, GABP, 53BP2, Lin12, glp-1, SW14, and SW16. The functions of the ankyrin repeats are compatible with a role in protein-protein interactions (Bork, Proteins (1993) 17(4):363; Lambert and Bennet, Eur. J. Biochem. (1993) 211:1; Kerr et al., Current Op. Cell Biol. (1992) 4:496; Bennet et al., J. Biol. Chem. (1980) 255:6424).

Basic Region Plus Leucine Zipper Transcription Factors (BZIP: Pfam Accession No. PF00170). SEQ ID NOS:108, 1714, 3931, and 4356 represent polynucleotides encoding novel members of the family of basic region plus leucine zipper transcription factors. The bZIP superfamily (Hurst, Protein Prof. [1995) 2:105; and Ellenberger, Curr. Opin. Struct. Biol. (1994) 4:12) of eukaryotic DNA-binding transcription factors encompasses proteins that contain a basic region mediating sequence-specific DNA-binding followed by a leucine zipper required for dimerization. The consensus pattern for this protein family is: [KR]-x(1,3)-[RKSAQ]-N-x(2)-[SAQ](2)-x-[RKTAENO]-x-R-x-[RK].

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DEAD and DEAH box families ATP-dependent helicases (Dead_box_helic, Pfam Accession No. PF00270, SEQ ID NOS:38, 415, and 5756 correspond to sequences encoding a polypeptides that are members of the DEAD box family. A number of eukaryotic and prokaryotic proteins have been characterized (Schmid et al., Mol. Microbiol. (1992) 6:283; Linder et al., Nature (1989) 337:121; Wassarman et al., Nature (1991) 349:463) on the basis of their structural similarity. All are involved in ATP-dependent, nucleic-acid unwinding. All DEAD box family members of the above proteins share a number of conserved sequence motifs, some of which are specific to the DEAD family while others are shared by other ATP-binding proteins or by proteins belonging to the helicases 'superfamily' (Hodgman, Nature (1988) 333:22 and Nature (1988) 333:578 (Errata)). One of these motifs, called the 'D-E-A-D-box', represents a special version of the B motif of ATP-binding proteins. Some other proteins belong to a subfamily which have His instead of the second Asp and are thus said to be 'D-E-A-H-box' proteins (Wassarman et al., Nature (1991) 349:463; Harosh et al., Nucleic Acids Res. (1991) 19-6331; Koonin et al., J. Gen. Virol. (1992) 73:989). The following signature patterns are used to identify member for both subfamilies: 1) [LIVMFI(2)-D-E-A-D-[RKEN]-x-[LIVMFI(3)-D-E-IALIVI-H-INECRI.

Defensins (defensins: Pfam Accession No. PF00323, SEQ ID NO:486 corresponds to a sequence encoding a polypeptide that is a member of the mammalian defensin family. Defensins are a family of structurally related cysteine-rich peptides which are active against many Gram-positive bacteria, fungi, and enveloped viruses (Lehrer et al., ASM News (1990) 56:315-318; Lehrer et al., Cell (1991) 64:229-230; Kagan et al., Toxicology (1994) 87:131-149; Lehrer et al., Annu. Rev. Immunol. (1993) 11:105-128; White et al., Curr. Opin. Struc/ Biol. (1995) 5:521-527). Some defensins inhibit to infection and neoplasia. The peptides known to belong to this family range in length from 29 to 35 amino acids and have seven invariant residues, including six cysteines that are all involved in intrachain disulfide bonds. The following consensus pattern is used to identify members of this protein family: C-x-C-x(3.5)-C-x(7)-G-x-C-x(9)-C-C.

EF Hand (Efhand: Pfam Accession No. PF00036). SEQ ID NO.4373 corresponds to a polynucleotide encoding a member of the EF-hand protein family, a calcium binding domain shared by many calcium-binding proteins belonging to the same evolutionary family (Kawasaki et al., Protein. Prof. (1995) 2:305-490). The domain is a twelve residue loop flanked on both sides by a twelve residue alpha-helical domain, with a calcium ion coordinated in a pentagonal bipyramidal configuration. The six residues involved in the binding are in positions 1, 3, 5, 7, 9 and 12; these residues are denoted by X, Y, Z, -Y, -X and -Z. The invariant Glu or Asp at position 12 provides two oxygens for liganding Ca (bidentate ligand). The consensus pattern includes the complete EF-hand loop as well as the first residue which follows the loop and which seem to always be hydrophobic: D-x-[DNS]-{ILIVFYW}-[DENSTG]-[DNQGHRK]-{GP}-[LIVMC]-[DENQSTAGC]-x(2)-[DE]-[LIVYWFYW].

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Epidermal Growth Factor (EGF: Pfam Accession No. PF00008), SEQ ID NO:3689 represents a polymucleotide encoding a member of the EGF family of proteins. The distinguishing characteristic of this family is the presence of a sequence of about thirty to forty amino acid residues found in epidermal growth factor (EGF) which has been shown to be present, in a more or less conserved form, in a large number of other proteins (Davis, New Biol. (1990) 2:410-419; Blomquist et al., Proc. Natl. Acad. Sci. U.S.A. (1984) 81:7363-7367; Barkert et al., Protein Nucl. Acid Erz. (1986) 29:54-86; Doolittle et al., Nature. (1984) 307:558-560; Appella et al., FEBS Lett. (1988) 231:1-4; Campbell and Bork, Curr. Opin. Struct. Biol. (1993) 3:385-392). A common feature of the domain is that the conserved pattern is generally found in the extracellular domain of membrane-bound proteins or in proteins known to be secreted. The EGF domain includes six cysteine residues which have been shown to be involved in disulfide bonds. The main structure is a two-stranded beta-sheet followed by a loop to a C-terminal short two-stranded sheet. Subdomains between the conserved cysteines strongly vary in length. The following consensus patterns are used to identify members of this family: C-x-C-x(5)-G-x(2)-C and C-x-C-x(5)-[GPI-FFWI-x(4,8)-C.

Ets Domain (Ets_Cterm: Pfam Accession No. PF00178). SEQ ID NO:6 represents a polynucleotide encoding a polypeptide with C-terminal homology in the ETS domain. Proteins of this family contain a conserved domain, the "ETS-domain," that is involved in DNA binding. The domain appears to recognize purine-rich sequences; it is about 85 to 90 amino acids in length, and is rich in aromatic and positively charged residues (Wasylyk et al., Eur. J. Biochem. (1993) 211:718). The ets gene family encodes a novel class of DNA-binding proteins, each of which binds a specific DNA sequence and comprises an ets domain that specifically interacts with sequences containing the common core tri-nucleotide sequence GGA. In addition to an ets domain, native ets proteins comprise other sequences which can modulate the biological specificity of the protein. Ets genes and proteins

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are involved in a variety of essential biological processes including cell growth, differentiation and development, and three members are implicated in oncogenic process.

Pleckstrin Homology (PH; Pfam Accession No. PF00169), SEQ ID NOS:228 and 6001 correspond to polynucleotides encoding members of the PH family. The pleckstrin homology domain is a domain of about 100 residues that occurs in a wide range of proteins involved in intracellular signaling or as constituents of the cytoskeleton (Mayer et al., Cell (1993) 73:629-630; Haslam et al., Nature (1993) 363:309-310; Musacchio et al., Trends Biochem. Sci. (1993) 18:343-348; Gibson et al., Trends Biochem. Sci. (1994) 19:349-353; Pawson, Nature (1995) 373:573-80; Ingley and Hemmings, J. Cell. Biochem. (1994) 56:436-443; Saraste and Hyvonen, Curr. Optn. Struct. Biol. (1995) 5:403-408). All known structures of PH domains have two perpendicular anti-parallel beta sheets, followed by a C-terminal amphipathic helix. The loops connecting the beta-strands differ greatly in length, making the PH domain relatively difficult to detect (Riddihough, Nat. Struct. Biol. (1994) 1:755-757). There are no totally invariant residues within this domain.

Protein Kinase (protkinase: Pfam Accession No. PF00069). SEQ ID NOS-9, 39, 69, 118, 229 and 4151 represent polynucleotides encoding protein kinases, which catalyze phosphorylation of proteins in a variety of pathways, and are implicated in cancer. Eukaryotic protein kinases (Hanks et al., FASEB J. (1995) 9:576; Hunter, Meth. Enzymol. (1991) 200:3; Hanks et al., Meth. Enzymol. (1991) 200:38; Hanks, Curv. Optn. Struct. Biol. (1991) 2:369; Hanks et al., Science (1988) 241:42) belong to a very extensive family of proteins that share a conserved catalytic core common to both serine/threonine and tyrosine protein kinases. There are a number of conserved regions in the catalytic domain of protein kinases. The first region, located in the N-terminal extremity of the catalytic domain, is a glycine-rich stretch of residues in the vicinity of a lysine residue, which has been shown to be involved in ATP binding. The second region, located in the central part of the catalytic domain, contains a conserved aspartic acid residue that is important for the catalytic activity of the enzyme (Knighton et al., Science (1991) 253:407).

The protein kinase profile includes two signature patterns for this second region: one specific for serine/threonine kinases and the other for tyrosine kinases. A third profile is based on the alignment in Hanks et al. [FASEB J. (1995) 9-576) and covers the entire catalytic domain. The consensus patterns are as follows: 1) [LIV]-G-{P}-G-{P}-FWMGSTNH]-[SGA]-{PW}-[LIVCAT]-{PD}-x-[GSTACLIVMFY]-x(5,18)-[LIVMFYWCSTAR]-[AIVP]-[LIVMFAGCKR]-K, where K binds ATP; 2) [LIVMFYC]-x-[HY]-x-D-[LIVMFY]-K-x(2)-N-[LIVMFYCT](3), where D is an active site residue; and 3) [LIVMFYC]-x-[HY]-x-D-[LIVMFY]-[RSTAC]-x(2)-N-[LIVMFYC], where D is an active site residue;

Retroviral aspartyl protease (RVP, Pfam Accession No. PF00077), SEQ ID NO:2038 corresponds to a polynucleotide encoding a member of the Aspartyl Protease family. Aspartyl proteases, also know as acid proteases, are a widely distributed family of proteolytic enzymes known to exist in vertebrates, fungi, plants, retroviruses and some plant viruses (Foltmann, Essays Biochem. (1981) 17:52-84; Davies, Annu. Rev. Biophys. Chem. (1990) 19:189-215; Rao et al., Biochemistry (1991) 30:4663-4671). Most retroviral aspartyl proteases (RVP) are homodimers of a chain of about 95 to 125 amino acids. In most retroviruses, the protease is encoded as a segment of a polyprotein which is cleaved during the maturation process of the virus. RVP is generally part of the pol polyprotein and, more rarely, of the gag polyprotein. The consensus pattern is as follows:

[LIVMFGAC]-[LIVMTADN]-[LIVFSA]-D-[STI-G-[STAV]-[STAPDENQ]-x-[LIVMFSTNC]-x[LIVMFGTAT] [D is the active site residue].

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Reverse Transcriptase (rvt. Pfam Accession No. PF00078). SEQ ID NOS:1511 and 2514 represent polynucleotides encoding reverse transcriptases, which occur in a variety of mobile elements, including retrotransposons, retroviruses, group II introns, bacterial msDNAs, hepadnaviruses, and caulimoviruses (Xiong and Eickbush, EMBOJ (1990) 9:3353-3362). Reverse transcriptases catalyze RNA-template-directed extension of the 3^3 -end of a DNA strand by one deoxynucleotide at a time and require an RNA or DNA primer.

Transforming growth factor beta like domain (TGF_beta: Pfam Accession No. PF00019).

SEQ ID NO.5522 represents a polynucleotide encoding a polypeptide of the TGF-beta family.

Proteins from the transforming growth factor-beta family are active as home- or hetero-dimers with the two chains being linked by a single disulfide bond (Roberts and Sporn, In Peptide growth factors and their receptors, Handbook of Experimental Pharmacology, Vol. 95, pp419-475, Springer Verlag, Heidelberg, (1990); Burt, Biochem. Biophys. Res. Commun. (1992) 184:590-595; Burt and Law, Prog. Growth Factor Res. (1994) 5:99-118). It is known from X-ray studies that all the other cysteines of the sequence are involved in interchain disulfide bonds (Daopin et al., Science (1992) 257:369-373). Members of this family can be recognized by the following consensus pattern: [LIVM]-x(2)-P-x(2)-[FY]-x(4)-C-x-G-x-C (The two C's are involved in disulfide bonds).

WD Domain (WD40), G-Beta Repeats (WD_domain; Pfam Accession No. PF00400). SEQ ID NO:117 represents a member of the WD domain/G-beta repeat family. Beta-transducin (G-beta) is one of the three subunits (alpha, beta, and gamma) of the guanine nucleotide-binding proteins (G proteins) which act as intermediaries in the transduction of signals generated by transmembrane receptors (Gilman, Annu. Rev. Blochem. (1987) 56:615). The alpha subunit binds to and hydrolyzes GTP; the beta and gamma subunits are required for the replacement of GDP by GTP as well as for membrane anchoring and receptor recognition. In higher eukaryotes, G-beta exists as a small

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multigene family of highly conserved proteins of about 340 amino acid residues. Structurally, G-beta has eight tandem repeats of about 40 residues, each containing a central Trp-Asp motif (this type of repeat is sometimes called a WD-40 repeat). The consensus pattern for the WD domain/G-Beta repeat family is: [LIVMSTAC]-[LIVMFYWSTAGC]-[LIMSTAG]-[LIVMSTAG]-[LIVMSTAG]-[LIVMFYSTAGC]-[LIVMSTAG]-[LIVMFSTAGC]-[LIVMSTAG]-[LIVMFSTAGC]-[LIVMST

Zinc Finger, C2H2 Type (Zincfing, C2H2; Pfam Accession No. PF00096). SEQ ID NOS; 61, 502, 700, 847, 2034, 2054, 3403, 3524, 3653, 3723, 4688, and 4979 correspond to polymucleotides encoding members of the C2H2 type zinc finger protein family, which contain zinc finger domains that facilitate nucleic acid binding (Klug et al., Trands Biochem. Scl. (1987) 12:464; Evans et al., Cell (1988) 52:1; Payre et al., FEBS Lett. (1988) 234:245; Miller et al., EMBO J. (1985) 4:1609; and Berg, Proc. Natl. Acad. Scl. USA (1988) 85:99). In addition to the conserved zinc ligand residues, a number of other positions are also important for the structural integrity of the C2H2 zinc fingers (Rossenfeld et al., J. Biomol. Struct. Dyn. (1993) 11:557). The best conserved position, which is generally an aromatic or aliphatic residue, is located four residues after the second cysteine. The consensus pattern for C2H2 zinc fingers is: C-x(2,4)-C-x(3)-[LIVMFYWC]-x(8)-H-x(3,5)-H. The two C's and two It's are zinc ligands.

Zine Finger. C2HC Type (Zinefing_C2HC: Pfam Accession No. PF01530). SEQ ID NO.3814 represents a member of the C2HC zine finger family. The 18 residue C2HC domain is mainly found in the retroviral nucleocapsid protein, and is required for viral genome packaging and for the early infection process (Katz and Jentoft, Bloessays (1989) 11:176-181; Urbaneja et al., JMol Blol. (1999) 287:59-75). In addition, the CCHC domain is found in eukaryotic proteins involved in RNA binding or single-strand DNA binding.

Zinc finger, C3HC4 type (RING finger), signature (Zincfing C3HC4: Pfam Accession No. PF00097). SEQ ID NO:31 40 represents a polymucleotide encoding a polypeptide having a 25 C3HC4 type zinc finger signature. A number of eukaryotic and viral proteins contain this signature, which is primarily a conserved cysteine-rich domain of 40 to 60 residues (Borden et al., Curr. Optn. Struct. Btol. (1996) 6:395) that binds two atoms of zinc, and is probably involved in mediating protein-protein interactions. The 3D structure of the zine ligation system is unique to the RING domain and is referred to as the "cross-brace" motif. The spacing of the cysteines in such a domain is Cxx(2)-Cx(9 to 39)-Cxx(1 to 3)-H-x(2 to 3)-C-x(2)-Cx(4 to 48)-C-x(2)-C. The signature pattern for the C3HC4 finger is based on the domain's central region: C-x-H-x-[LIVMFY]-C-x(2)-C-[LIVMYA].

Zinc finger, CCCH type, (Zincfing, CCCH; Pfam Accession No. PF00642). SEQ ID NO:877 corresponds to a polyncleotide encoding a member of the CCCH zinc finger protein family. This domain is present in many eukaryotic proteins, including zinc finger proteins involved in cell cycle or

growth phase-related regulation and regulatory proteins involved in regulating the response to growth factors. It has been shown that proteins containing the CCCH zinc finger interact with the 3' untranslated region of various mRNA (Carballo et al., Science (1998) 281:1001-1005; Lai et al., Mol. Cell. Biol. (1999) 19:4311-4323) and that this domain is often present in two copies. The consensus pattern for the CCCH zinc fingers is: C-x8-C-x5-C-x3-H.

Example 4: Description of Libraries and Detection of Differential Expression

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The relative expression levels of the polynucleotides of the invention were assessed in several libraries prepared from various sources, including cell lines and patient tissue samples. Table 1 above provides a summary of these libraries, including the shortened library name, the mRNA source used to prepared the cDNA library, the "nickname" of the library that is used in the tables below (in quotes). and the approximate number of clones in the library.

Each of the libraries is composed of a collection of cDNA clones that in turn are representative of the mRNAs expressed in the indicated mRNA source. In order to facilitate the analysis of the millions of sequences in each library, the sequences were assigned to clusters. The concept of "cluster of clones" is derived from a sorting/grouping of cDNA clones based on their hybridization pattern to a panel of roughly 300 7bp oligonucleotide probes (see Drmanac et al., Genomics (1996) 37(1):29). Random cDNA clones from a tissue library are hybridized at moderate stringency to 300 7bp oligonucleotides. Each oligonucleotide has some measure of specific hybridization to that specific 20 clone. The combination of 300 of these measures of hybridization for 300 probes equals the "hybridization signature" for a specific clone. Clones with similar sequence will have similar hybridization signatures. By developing a sorting/grouping algorithm to analyze these signatures. groups of clones in a library can be identified and brought together computationally. These groups of clones are termed "clusters". Depending on the stringency of the selection in the algorithm (similar to the stringency of hybridization in a classic library cDNA screening protocol), the "purity" of each cluster can be controlled. For example, artifacts of clustering may occur in computational clustering just as artifacts can occur in "wet-lab" screening of a cDNA library with 400 bp cDNA fragments, at even the highest stringency. The stringency used in the implementation of cluster herein provides groups of clones that are in general from the same cDNA or closely related cDNAs. Closely related clones can be a result of different length clones of the same cDNA, closely related clones from highly related gene families, or splice variants of the same cDNA.

Differential expression for a selected cluster was assessed by first determining the number of cDNA clones corresponding to the selected cluster in the first library (Clones in 1th), and the determining the number of cDNA clones corresponding to the selected cluster in the second library

(Clones in 2nd). Differential expression of the selected cluster in the first library relative to the second library is expressed as a "ratio" of percent expression between the two libraries. In general, the "ratio" is calculated by: 1) calculating the percent expression of the selected cluster in the first library by dividing the number of clones corresponding to a selected cluster in the first library by the total number of clones analyzed from the first library; 2) calculating the percent expression of the selected cluster in the second library by dividing the number of clones corresponding to a selected cluster in a second library by the total number of clones analyzed from the second library; 3) dividing the calculated percent expression from the first library by the calculated percent expression from the first library by the calculated percent expression from the second library. If the "number of clones" corresponding to a selected cluster in a library is zero, the value is set at 1 to aid in calculation. The formula used in calculating the ratio takes into account the "depth" of each of the libraries being compared, i.e., the total number of clones analyzed in each library.

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In general, a polynucleotide is significantly differentially expressed between two samples when the ratio value is greater than at least about 2, preferably greater than at least about 3, more preferably greater than at least about 5, where the ratio value is calculated using the method described above. The significance of differential expression is determined using a z score test (Zar, <u>Biostatistical Analysis</u>, Prentice Hall, Inc., USA, "Differences Between Proportions," pp 296-298 (1974).

Using this approach, a number of polymucleotide sequences were identified as being differentially expressed between, for example, cells derived from high metastatic potential cancer tissue and low metastatic cancer cells, and between cells derived from metastatic cancer tissue and normal tissue. Evaluation of the levels of expression of the genes corresponding to these sequences can be valuable in diagnosis, prognosis, and/or treatment (e.g., to facilitate rationale design of therapy, monitoring during and after therapy, etc.). Moreover, the genes corresponding to differentially expressed sequences described herein can be therapeutic targets due to their involvement in regulation (e.g., inhibition or promotion) of development of, for example, the metastatic phenotype. For example, sequences that correspond to genes that are increased in expression in high metastatic potential cells relative to normal or non-metastatic tumor cells may encode genes or regulatory sequences involved in processes such as angiogenesis, differentiation, cell replication, and metastasis.

Detection of the relative expression levels of differentially expressed polynucleotides described herein can provide valuable information to guide the clinician in the choice of therapy. For example, a patient sample exhibiting an expression level of one or more of these polynucleotides that corresponds to a gene that is increased in expression in metastatic or high metastatic potential cells may warrant more aggressive treatment for the patient. In contrast, detection of expression levels of a polynucleotide sequence that corresponds to expression levels associated with that of low metastatic potential cells may warrant a more positive prognosis than the gross pathology would suggest.

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A number of polynucleotide sequences of the present invention are differentially expressed between human microvascular endothelial cells (HMVEC) that have been treated with growth factors relative to untreated HMVEC. Sequences that are differentially expressed between growth factor-treated HMVEC. Sequences that are differentially expressed between growth factor-treated HMVEC and untreated HMVEC can represent sequences encoding gene processes. For example, sequences that are more highly expressed in HMVEC treated with growth factors (such as bFGF or VEGF) relative to untreated HMVEC can serve as drug targets for chemotherapeutics, e.g., decreasing expression of such up-regulated genes or inhibiting the activity of the encoded gene product would serve to inhibit tumor cell angiogenesis. Detection of expression of these sequences in colon cancer tissue can be valuable in determining diagnostic, prognostic and/or treatment information associated with the prevention of achieving the malignant state in these tissues, and can be important in risk assessment for a patient. A patient sample displaying an increased level of one or more of these polynucleotides may thus warrant closer attention or more frequent screening procedures to catch the malignant state as early as possible.

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The differential expression of the polynucleotides can thus be used as, for example, diagnostic and/or prognostic markers, for risk assessment, patient treatment and the like. These polynucleotides can also be used in combination with other molecular and/or biochemical markers.

The differential expression data for polynucleotides of the invention that have been identified as being differentially expressed across various combinations of the libraries described above is summarized in Table 5 (inserted prior to the claims). Table 5 provides: 1) the Sequence Identification Number ("SEQ ID NO") assigned to the polynucleotide; 2) the cluster ("CLUSTER") to which the polynucleotide has been assigned as described above; 3) the library comparisons that resulted in identification of the polynucleotide as being differentially expressed ("PAIR AB"), where the cDNA libraries used are referenced by their library numbers; 4) the number of clones corresponding to the polynucleotide in the first library listed ("CLONES A"); 5) the number of clones corresponding to the polynucleotide in the second library listed ("CLONES B"); 6) the "RATIO PLUS" where the comparison resulted in a finding that the number of clones in library A is greater than the number of clones in library B is greater than the number of clones in library A.

Detection of expression of genes that correspond to the above polynucleotides may be of particular interest in diagnosis, prognosis, risk assessment, and monitoring of treatment. Furthermore, differential expression of a specific gene across multiple libraries can also be indicative of a gene whose expression is associated with, for example, suppression of the metastatic phenotype or with development of the cell toward a metastatic phenotyne. For example, SEO ID NO:3744 corresponds

to a gene that is expressed at relatively higher levels in metastatized colon tumor than in normal colon tissue. Thus a relatively increased level of expression of the gene corresponding to SEQ ID NO:3744 may be used as marker of a metastatic or pre-metastatic colon cels either alone or in combination with other markers.

Some polynucleotides exhibited similar differential expression trends in libraries of different tissue origin (see, e.g., SEQ ID NO:337). These data suggest that the differential expression patterns of some genes associated with development of tumors indicate a role for those genes that is nonspecific to the tissue of origin.

10 Example 5: Detection of Differential Expression Using Arrays

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mRNA isolated from samples of cancerous and normal colon tissue obtained from patients were analyzed to identify genes differentially expressed in cancerous and normal cells. Normal and cancerous cells collected from cryopreserved patient tissues were isolated using laser capture microdissection (LCM) techniques, which techniques are well known in the art (see, e.g., Ohyama et al. (2000) Biotechniques 29:530-6; Curran et al. (2000) Mol. Pathol. 53:64-8; Suarcz-Quian et al. (1999) Biotechniques 26:328-35; Simone et al. (1998) Trends Genet 14:272-6; Conia et al. (1997) J. Clin. Lab. Anal. 11:28-38; Emmert-Buck et al. (1996) Science 274:998-1001).

Table 6 (inserted before the claims) provides information about each patient from which colon tissue samples were isolated, including: the Patient ID ("PT ID")and Path ReportID (}Path ID"), which are numbers assigned to the patient and the pathology reports for identification purposes; the group ("Grp")to which the patients have been assigned; the anatomical location of the tumor ("Anatom Loc"); the primary tumor size ("Size"); the primary tumor grade ("Grade"); the identification of the histopathological grade ("Histo Grade"); a description of local sites to which the tumor had invaded ("Local Invasion"); the presence of lymph node metastases ("LN Met"); the incidence of lymph node metastases (provided as a number of lymph nodes positive for metastasis over the number of lymph nodes examined) ("Incidence Lymphnode Met"); the "Regional Lymphnode Grade"; the identification or detection of metastases to sites distant to the tumor and their location ("Dist Met & Loc"); the grade of distant metastasis ("Dist Met Grade"); and general comments about the patient or the tumor ("Comments"). Histophatology of all primary tumors incidated the tumor was adenocarcinmoa except for Patient ID Nos. 130 (for which no information was provided), 392 (in which greater than 50% of the cells were mucinous carcinoma), and 784 (adenosquamous carcinoma). Extranodal extensions were described in three patients, Patient ID Nos. 784, 789, and 791. Lymphovascular invasion was described in Patient ID Nos. 128, 278, 517, 534, 784, 786, 789, 791, 890, and 892. Crohn's-like

infiltrates were described in seven patients, Patient ID Nos. 52, 264, 268, 392, 393, 784, and 791.

Table 7 (below) provides information about the patients from whom the prostate tissue was isolated.

Table 7. Prostate paitent data.

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Prostate Patient ID		Normal Prostate Description		
96	3+3 Adenocarcinoma	Normal prostate; Benign hyperplasia		
282	4+3 Adenocarcinoma	Normal prostate; Benign hyperplasia		
286	3+3 Adenocarcinoma	Normal prostate; Benign hyperplasia		
294	3+4 Adenocarcinoma	Normal prostate; Benign hyperplasia		
362		Normal prostate; Benign hyperplasia		
428	4+3 Adenocarcinoma	Normal prostate; Benign hyperplasia		
492	3+3 Adenocarcinoma	Normal prostate; Benign hyperplasia		
492	3+3 Adenocarcinoma	Normal prostate; Benign hyperplasia		
493	3+4 Adenocarcinoma	Normal prostate; Benign hyperplasia		
510	3+3 Adenocarcinoma	Normal Prostate; Benign hyperplasia		

Identification of differentially expressed genes

cDNA probes were prepared from total RNA isolated from the patient cells described above.

Since LCM provides for the isolation of specific cell types to provide a substantially homogenous cell sample, this provided for a similarly pure RNA sample.

Total RNA was first reverse transcribed into cDNA using a primer containing a T7 RNA polymerase promoter, followed by second strand DNA synthesis. cDNA was then transcribed in vitro to produce antisense RNA using the T7 promoter-mediated expression (see, e.g., Luo et al. (1999) Nature Med 5:117-122), and the antisense RNA was then converted into cDNA. The second set of cDNAs were again transcribed in vitro, using the T7 promoter, to provide antisense RNA. Optionally, the RNA was again converted into cDNA, allowing for up to a third round of T7-mediated amplification to produce more antisense RNA. Thus the procedure provided for two or three rounds of in vitro transcription to produce the final RNA used for fluorescent labeling.

Fluorescent probes were generated by first adding control RNA to the antisense RNA mix, and producing fluorescently labeled cDNA from the RNA starting material. Fluorescently labeled cDNAs prepared from the tumor RNA sample were compared to fluorescently labeled cDNAs prepared from normal cell RNA sample. For example, the cDNA probes from the normal cells were labeled with Cy3 fluorescent dye (green) and the cDNA probes prepared from the tumor cells were labeled with Cy5 fluorescent dye (red), and vice versa.

Each array used had an identical spatial layout and control spot set. Each microarray was divided into two areas, each area having an array with, on each half, twelve groupings of 32 x 12 spots, for a total of about 9,216 spots on each array. The two areas are spotted identically which provide for at least two duplicates of each clone per array.

Polynucleotides for use on the arrays were obtained from both publicly available sources and from cDNA libraries generated from selected cell lines and patient tissues. PCR products of from about 0.5kb to 2.0 kb amplified from these sources were spotted onto the array using a Molecular Dynamics Gen III spotter according to the manufacturer's recommendations. For polynucleotides described herein, the microarray spot contained a clone having a cDNA from which the sequence was derived. The first row of each of the 24 regions on the array had about 32 control spots, including 4 negative control spots and 8 test polynucleotides. The test polynucleotides were spiked into each sample before the labeling reaction with a range of concentrations from 2-600 pg/slide and ratios of 1:1. For each array design, two slides were hybridized with the test samples reverse-labeled in the labeling reaction. This provided for about four duplicate measurements for each clone, two of one color and two of the other, for each sample.

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Table 8 (inserted before the claims) describes sequences present on the arrays. Table 8 includes: 1) athe SEQ ID NO of the sequence of the polynucleotide; and 2)the Spot ID, which is a unique identifier for each spot ontaining target sequence of interest on all arrays used.

The differential expression assay was performed by mixing equal amounts of probes from tumor cells and normal cells of the same patient. The arrays were prehybridized by incubation for about 2 hrs at 60°C in 5X SSC/0.2% SDS/1 mM EDTA, and then washed three times in water and twice in isopropanol. Following prehybridization of the array, the probe mixture was then hybridized to the array under conditions of high stringency (overnight at 42°C in 50% formamide, 5X SSC, and 0.2% SDS. After hybridization, the array was washed at 55°C three times as follows: 1) first wash in 1X SSC/0.2% SDS; and 3) third wash in 0.1X SSC.

The arrays were then scanned for green and red fluorescence using a Molecular Dynamics Generation III dual color laser-scanner/detector. The images were processed using BioDiscovery Autogene software, and the data from each scan set normalized to provide for a ratio of expression relative to normal. Data from the microarray experiments was analyzed according to the algorithms described in U.S. application scrial no. 60/252,358, filed November 20, 2000, by E.J. Moler, M.A. Boyle, and F.M. Randazzo, and entitled "Precision and accuracy in cDNA microarray data," which application is specifically incorporated herein by reference.

The experiment was repeated, this time labeling the two probes with the opposite color in order to perform the assay in both "color directions." Each experiment was sometimes repeated with two more slides (one in each color direction). The level fluorescence for each sequence on the array expressed as a ratio of the geometric mean of 8 replicate spots/genes from the four arrays or 4 replicate spots/gene from 2 arrays or some other permutation. The data were normalized using the spiked positive controls present in each duplicated area, and the precision of this normalization was included

in the final determination of the significance of each differential. The fluorescent intensity of each spot was also compared to the negative controls in each duplicated area to determine which spots have detected significant expression levels in each sample.

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A statistical analysis of the fluorescent intensities was applied to each set of duplicate spots to assess the precision and significance of each differential measurement, resulting in a p-value testing the null hypothesis that there is no differential in the expression level between the tumor and normal samples of each patient. During initial analysis of the microarrays, the hypothesis was accepted if p> 10⁻³, and the differential ratio was set to 1.000 for those spots. All other spots have a significant difference in expression between the tumor and normal sample. If the tumor sample has detectable expression and the normal does not, the ratio is truncated at 1000 since the value for expression in the normal sample would be zero, and the ratio would not be a mathematically useful value (e.g., infinity). If the normal sample has detectable expression and the tumor does not, the ratio is truncated to 0.001. since the value for expression in the tumor sample would be zero and the ratio would not be a mathematically useful value. These latter two situations are referred to herein as "on/off." Database tables were populated using a 95% confidence level (p>0.05).

Table 8 (inserted before the claims) provides the results for gene products differentially expressed in the colon tumor samples relative to normal tissue samples. Table 8 includes: 1) the SEQ ID NO; 2) the spot identification number ("SpotID"); 3) the percentage of patients tested in which expression levels of the gene (as detected using the correponding clone) was at least 2-fold greater in cancerous colon tissue (primary colon tumor) than in matched normal tissue ("Colon>2x T/N"); 4) the percentage of patients tested in which expression levels of the gene was less than or equal to one-half of the expression level in matched normal cells ("Colon <=halfx T/N"); and 5) the colon number ratios. indicating the number of patients upon which the provided ratios was based.

Table 9 below provides the data for differential expression analysis on the arrays using samples from metastazed colon tissue. In this example, the samples used for hybridization to sequences on the microarray were derived from the matched metastasized (MT) colon tissue and normal (N) colon tissues of the patients. Table 9 includes: 1) the SEQ ID NO; 2) the percentage of patients tested in which expression levels of the gene (as detected using the correponding clone) was at least 2-fold greater in metastisized cancerous colon tissue (MT) than in matched normal tissue 30 ("Colon>2x MT/N"); 5) the percentage of patients tested in which expression levels of the gene was less than or equal to one-half of the expression level in matched normal cells ("Colon <=halfx T/N"); and 8) the colon number ratios, indicating the number of patients upon which the provided ratios was based. The corresponding data with the same sequence of the colon tumor tissue versus matched normal colon tissue (T/N) are provided for convenience in comparison,

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Table 9. Polynucleotides Corresponding to Differnetially Expressed Genes in Metastasized Colon Cancer Tissue

			Colon MT/N			
			Num Ratios	Colon T/N	Colon T/N	Colon T/N
SEQ ID NO	Colon MT/N >2x	Colon MT/N <halfx< td=""><td>by Clone</td><td>>2x</td><td><halfx< td=""><td>Num Ratios</td></halfx<></td></halfx<>	by Clone	>2x	<halfx< td=""><td>Num Ratios</td></halfx<>	Num Ratios
217	40.0	0.0	5.0	0.0	. 50.0	8.0
324	0.0	40.0	5.0	0.0	37,5	8.0
1653	0.0	40.0	5.0	0.0	37.5	8.0
1972	40.0	0.0	5.0	0.0	42.9	7.0
2346	20.0	40.0	5.0	0.0	85.7	7.0
2352	20.0	80.0	5.0	0.0	71.4	7.0
2353	20.0	_40.0	5.0	0.0	85.7	7.0
2647	0.0	40.0	5.0	0.0	62.5	8.0
5341	0.0	40.0	5.0	0.0	37.5	8.0

Table 10 below provides the data for differential expression analysis on the arrays using samples from matched cancerous and normal prostate tissue (PT/N). Table 10 includes: 1) the SEQ ID NO; 2) the percentage of patients tested in which expression levels of the gene (as detected using the corresponding clone) was at least 2-fold greater in metastisized cancerous prostate tissue (PT) than in matched normal tissue ("Colon>2x PT/N"); 3) the percentage of patients tested in which expression levels of the gene was less than or equal to one-half of the expression level in matched normal cells ("Colon <=halfx PT/N"); and 4) the prostate PT/N number ratios, indicating the number of patients upon which the provided ratios was based. The corresponding data with the same sequences for the colon tumor versus normal (T/N) and metastasized colon tissue versus normal (MT/N) are provided for convenience in comparison.

Table 10. Polynucleotides Corresponding to Differnetially Expressed Genes in Prostate

Cancer Tissue									
SEQ ID NO	Prostate (PT/N) >2x	Prostate (PT/N) <halfx< td=""><td>Prostate (PT/N) Num Ratios</td><td>Colon T/N >2x</td><td>Colon T/N <halfx< td=""><td>Colon T/N Num Ratios</td><td>Colon MT/N >2x</td><td>Colon MT/N <halfx< td=""><td>Colon MT/N Num Ratios</td></halfx<></td></halfx<></td></halfx<>	Prostate (PT/N) Num Ratios	Colon T/N >2x	Colon T/N <halfx< td=""><td>Colon T/N Num Ratios</td><td>Colon MT/N >2x</td><td>Colon MT/N <halfx< td=""><td>Colon MT/N Num Ratios</td></halfx<></td></halfx<>	Colon T/N Num Ratios	Colon MT/N >2x	Colon MT/N <halfx< td=""><td>Colon MT/N Num Ratios</td></halfx<>	Colon MT/N Num Ratios
139	11.1	33.3	9.0	0.0	50.0	8.0			
490	37.5	12.5	8.0	0.0	71.4	7.0			
629	33,3	11.1	9.0						
644	12.5	37.5	8.0	0.0	42.9	7.0			
1674	33.3	0.0	9.0						
2346	37.5	25.0	8.0	0.0	85.7	7.0	20.0	40.0	5.0
2352	37.5	12.5	8.0	0.0	71.4	7.0	20.0	80.0	5.0
2420	22.2	33.3	9.0						

		Prostate (PT/N) <halfx< th=""><th></th><th>Colon T/N >2x 0.0</th><th>Colon T/N <halfx< th=""><th>Colon T/N Num Ratios 8.0</th><th></th><th>Colon MT/N <halfx< th=""><th>Num</th></halfx<></th></halfx<></th></halfx<>		Colon T/N >2x 0.0	Colon T/N <halfx< th=""><th>Colon T/N Num Ratios 8.0</th><th></th><th>Colon MT/N <halfx< th=""><th>Num</th></halfx<></th></halfx<>	Colon T/N Num Ratios 8.0		Colon MT/N <halfx< th=""><th>Num</th></halfx<>	Num
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Example 6: Antisense Regulation of Gene Expression

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The expression of the differentially expressed genes represented by the polynucleotides in the cancerous cells can be further analyzed using antisense knockout technology to confirm the role and function of the gene product in tumorigenesis, e, g,, in promoting a metastatic phenotype.

Methods for analysis using antisense technology are well known in the art. For example, a number of different oligonucleotides complementary to the mRNA generated by the differentially expressed genes identified herein can be designed as antisense oligonucleotides, and tested for their ability to suppress expression of the genes. Sets of antisense oligoners specific to each candidate target are designed using the sequences of the polynucleotides corresponding to a differentially expressed gene and the software program HYBsimulator Version 4 (available for Windows 95/Windows NT or for Power Macintosh, RNAture, Inc. 1003 Health Sciences Road, West, Irvine, CA 92612 USA). Factors considered when designing antisense oligonucleotides include: 1) the secondary structure of oligonucleotides; 2) the secondary structure of the target gene; 3) the specificity with no or minimum cross-hybridization to other expressed genes; 4) stability; 5) length and 6) terminal GC content. The antisense oligonucleotide is designed to so that it will hybridize to its target sequence under conditions of high stringency at physiological temperatures (e.g., an optimal temperature for the cells in culture to provide for hybridization in the cell, e.g., about 37°C), but with minimal formation of homodimers.

Once synthesized and quantitated, the oligomers are screened for efficiency of a transcript knock-out in a panel of cancer cell lines. The efficiency of the knock-out is determined by analyzing mRNA levels using lightcycler quantification. The oligomers that resulted in the highest level of transcript knock-out, wherein the level was at least about 50%, preferably about 80-90%, up to 95% or more up to undetectable message, are selected for use in a cell-based proliferation assay, an anchorage independent growth assay, and an apoptosis assay.

For example, where the polynucleotide is identified as having a role in colon cancer, the ability of the corresponding designed antisense oligonucleotide to inhibit gene expression is tested through transfection into SW620 colon colorectal carcinoma cells. For each transfection mixture, a carrier molecule, preferably a lipitoid or cholesteroid, is prepared to a working concentration of 0.5 mM in water, sonicated to yield a uniform solution, and filtered through a 0.45 µm PVDF membrane. The

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antisense or control oligonucleotide is then prepared to a working concentration of $100 \, \mu M$ in sterile Millipore water. The oligonucleotide is further diluted in OptiMEMTM (Giboo/BRL), in a microfuge tube, to $2 \, \mu M$, or approximately $20 \, \mu g$ oligo/ml of OptiMEMTM. In a separate microfuge tube, lipitoid or cholesteroid, typically in the amount of about $1.5\text{-}2 \, \text{nmol}$ lipitoid/ μg antisense oligonucleotide, is diluted into the same volume of OptiMEMTM used to dilute the oligonucleotide. The diluted antisense oligonucleotide is immediately added to the diluted lipitoid and mixed by pipetting up and down. Oligonucleotide is added to the cells to a final concentration of $30 \, n M$.

The level of target mRNA that corresponds to a target gene of interest in the transfected cells is quantitated in the cancer cell lines using the Roche LightCycler TM real-time PCR machine. Values for the target mRNA are normalized versus an internal control (e.g., beta-actin). For each $20~\mu$ l reaction, extracted RNA (generally 0.2-1 μ g total) is placed into a sterile $0.5~\sigma$ 1.5 ml microcentrifuge tube, and water added to a total volume of $12.5~\mu$ l. To each tube $7.5~\mu$ l of a buffer/enzyme mixture is added, which is prepared by mixing (in the order listed) $2.5~\mu$ l H_2O , $2.0~\mu$ l 10X reaction buffer, $10~\mu$ l oligo dT ($20~\mu$) mix ($10~\mu$ mix ($10~\mu$) meach), $0.5~\mu$ l RNAsin® ($20~\mu$) (Ambion, Inc., Hialeah, FL), and $0.5~\mu$ l MMLV reverse transcriptase ($50~\mu$) (Ambion, Inc.). The contents are mixed by pipetting up and down, and the reaction mixture incubated at $4.2~\mu$ C for $1~\mu$ hour. The contents of each tube are centrifuged prior to amplification.

An amplification mixture is prepared by mixing in the following order: LX PCR buffer II, 3
mM MgCl₂, 140 µM each dNTP, 0.175 pmol each oligo, 1:50,000 dil of SYBR® Green, 0.25 mg/ml

BSA, 1 unit Taq polymerase, and H₂O to 20 µl. (PCR buffer II is available in 10X concentration from

Perkin-Elmer, Norwalk, CT). In 1X concentration it contains 10 mM Tris pH 8.3 and 50 mM KCl.

SYBR® Green (Molecular Probes, Eugene, OR) is a dye which fluoresces when bound to double

stranded DNA. As double stranded PCR product is produced during amplification, the fluorescence

from SYBR® Green increases. To each 20 µl aliquot of amplification mixture, 2 µl of template RT are

added, and amplification carried out according to standard protocols.

The results can be expressed as the percent decrease in expression of the corresponding gene product relative to non-transfected cells, vehicle-only transfected (mock-transfected) cells, or cells transfected with reverse control oligonucleotides.

30 Example 7: Effect of Expression on Proliferation

The effect of gene expression on the inhibition of cell proliferation can be assessed in, for example, metastatic breast cancer cell lines (MDA-MB-231 ("231")), SW620 colon colorectal carcinoma cells, or SKOV3 cells (a human ovarian carcinoma cell line).

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Cells are plated to approximately 60-80% confluency in 96-well dishes. Antisense or reverse control oligonucleotide iss diluted to 2 µM in OptiNEM™ and added to OptiMEM™ into which the delivery vehicle, lipitoid 11:6-6 in the case of SW620 cells or 1:1 lipitoid 1:cholesteroid 1 in the case of MDA-MB-231 cells, had been diluted. The oligo/delivery vehicle mixture is then further diluted into medium with serum on the cells. The final concentration of oligonucleotide for all experiments was 300 nM, and the final ratio of oligo to delivery vehicle for all experiments iss 1.5 nmol lipitoid/µg oligonucleotide.

Antisense oligonucleotides are prepared as described above (see Example 6). Cells are transfected overnight at 37°C and the transfection mixture replaced with fresh medium the next morning. Transfection is carried out as described above in Example 3.

Those antisense oligonucleotides that inhibit proliferation represent genes that play a role in production or maintenance of the cancerous phenotype.

Example 8: Effect of Gene Expression on Colony Formation

The effect of gene expression upon colony formation of, for example, SW620 cells, SKOV3 cells, and MD-MBA-231 cells can be tested in a soft agar assay. Soft agar assays are conducted by first establishing a bottom layer of 2 ml of 0.6% agar in media plated fresh within a few hours of layering on the cells. The cell layer is formed on the bottom layer by removing cells transfected as described above from plates using 0.05% trypsin and washing twice in media. The cells are counted in a Coulter counter, and resuspended to 10^6 per ml in media. $10~\mu$ 1 aliquots are placed with media in 96-well plates (to check counting with WST1), or diluted further for the soft agar assay. 2000 cells are plated in 800 μ 1 0.4% agar in duplicate wells above 0.6% agar bottom layer. After the cell layer agar solidifies, 2 ml of media is dribbled on top and antisense or reverse control oligo (produced as described in Example 6) added without delivery vehicles. Fresh media and oligos are added every 3-4 days. Colonies usually are expected to form in 10 days to 3 weeks. Fields of colonies are counted by eye. Wst-1 metabolism values can be used to compensate for small differences in starting cell number. Larger fields can be seanned for visual record of differences.

Those antisense oligonucleotides that inhibited colony formation represent genes that play a role in production or maintenance of the cancerous phenotype.

Example 9: Induction of Cell Death upon Depletion of Polypeptides by Depletion of mRNA ("Antisense Knockout")

In order to assess the effect of depletion of a target message upon cell death, SW620 cells, or other cells derived from a cancer of interest, are transfected for proliferation assays. For cytotoxic

effect in the presence of cisplatin (cis), the same protocol is followed but cells are left in the presence of 2 μ M drug. Each day, cytotoxicity was monitored by measuring the amount of LDH enzyme released in the medium due to membrane damage. The activity of LDH is measured using the Cytotoxicity Detection Kit from Roche Molecular Biochemicals. The data is provided as a ratio of LDH released in the medium vs. the total LDH present in the well at the same time point and treatment (rLDH/tLDH). A positive control using antisense and reverse control oligonucleotides for BCL2 (a known anti-apoptotic gene) is included; loss of message for BCL2 leads to an increase in cell death compared with treatment with the control oligonucleotide (background cytotoxicity due to transfection).

Example 10: Functional Analysis of Gene Products Differentially Expressed in Cancer

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The gene products of sequences of a gene differentially expressed in cancerous cells can be further analyzed to confirm the role and function of the gene product in tumorigenesis, e.g., in promoting or inhibiting development of a metastatic phenotype. For example, the function of gene products corresponding to genes identified herein can be assessed by blocking function of the gene products in the cell. For example, where the gene product is secreted or associated with a cell surface membrane, blocking antibodies can be generated and added to cells to examine the effect upon the cell phenotype in the context of, for example, the transformation of the cell to a cancerous, particularly a metastatic, observive.

Where the gene product of the differentially expressed genes identified herein exhibits sequence homology to a protein of known function (e.g., to a specific kinase or protease) and/or to a protein family of known function (e.g., contains a domain or other consensus sequence present in a protease family or in a kinase family), then the role of the gene product in tumorigenesis, as well as the activity of the gene product, can be examined using small molecules that inhibit or enhance function of the corresponding protein or protein family.

Additional functional assays include, but are not necessarily limited to, those that analyze the effect of expression of the corresponding gene upon cell cycle and cell migration. Methods for performing such assays are well known in the art.

30 Example 11: Contig Assembly and Additional Gene Characterization

The sequences of the polynucleotides provided in the present invention can be used to extend the sequence information of the gene to which the polynucleotides correspond (e.g., a gene, or mRNA encoded by the gene, having a sequence of the polynucleotide described herein). This expanded sequence information can in turn be used to further characterize the corresponding gene, which in turn

provides additional information about the nature of the gene product (e.g., the normal function of the gene product). The additional information can serve to provide additional evidence of the gene product's use as a therapeutic target, and provide further guidance as to the types of agents that can modulate its activity.

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For example, a contig can be assembled using the sequence of a polynucleotide described herein. A "contig" is a contiguous sequence of nucleotides that is assembled from nucleic acid sequences having overlapping (e.g., shared or substantially similar) sequence information. The sequences of publicly-available ESTs (Expressed Sequence Tags) and the sequences of various clones from several cDNA libraries synthesized at Chiron were used in the contig assembly. The contig is assembled using the software program Sequencher, version 4.05, according to the manufacturer's instructions. The resulting contig can then be used to search both the public databases as well as databases internal to the applicatus to matchibe polynucleotide contiged with homology data and/or differential gene expressed data.

The sequence information obtained in the contig assembly described above can be used to obtain a consensus sequence derived from the contig using the Sequencher program. The consensus sequence can then be used as a query sequence in a BLASTN search of the DGTI DoubleTwist Gene Index (DoubleTwist, Inc., Oakland, CA), which contains all the EST and non-redundant sequence in public databases. Alternatively, a sequence of a polynucleotide described herein can be used directly as a query sequence in a BLASTN search of the DGTI DoubleTwist Gene Index.

Through contig assembly and the use of homology searching software programs, the sequence information provided herein can be readily extended to confirm, or confirm a predicted, gene having the sequence of the polynucleotides described in the present invention. Further the information obtained can be used to identify the function of the gene product of the gene corresponding to the polynucleotides described herein. While not necessary to the practice of the invention, identification of the function of the corresponding gene, can provide guidance in the design of therapeutics that target the gene to modulate its activity and modulate the cancerous phenotype (e.g., inhibit metastasis, proliferation, and the like).

Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it is readily apparent to those of ordinary skill in the art in light of the teachings of this invention that certain changes and modifications may be made thereto without departing from the spirit or scope of the appended claims. Those skilled in the art will recognize, or be able to ascertain, using not more than routine experimentation, many equivalents to the

specific embodiments of the invention described herein. Such specific embodiments and equivalents are intended to be encompassed by the following claims.

Deposit Information.

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A deposit of the biological materials in the tables referenced below was made with the American Type Culture Collection, 10801 University Blvd., Manasas, VA 20110-2209, under the provisions of the Budapest Treaty, on or before the filing date of the present application. The accession number indicated is assigned after successful viability testing, and the requisite fees were paid. Access to said cultures will be available during pendency of the patent application to one determined by the Commissioner to be entitled to such under 37 C.F.R. §1.14 and 35 U.S.C. §122. All restriction or availability of said cultures to the public will be irrevocably removed upon the granting of a patent based upon the application. Moreover, the designated deposits will be maintained for a period of thirty (30) years from the date of deposit, or for five (5) years after the last request for the deposit; or for the enforceable life of the U.S. patent, whichever is longer. Should a culture become nonviable or be inadvertently destroyed, or, in the case of plasmid-containing strains, lose its plasmid, it will be replaced with a viable culture(s) of the same taxonomic description.

These deposits are provided merely as a convenience to those of skill in the art, and are not an admission that a deposit is required. A license may be required to make, use, or sell the deposited materials, and no such license is hereby granted. The deposit below was received by the ATCC on or before the filing date of the present application.

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Table 11. Cell Lines Deposited with ATCC

Cell Line	Deposit Date	ATCC Accession No.	CMCC Accession No.		
KM12L4-A	March 19, 1998	CRL-12496	11606		
Km12C	May 15, 1998	CRL-12533	11611		
MDA-MB-	May 15, 1998	CRL-12532	10583		
231					
MCF-7	October 9, 1998	CRL-12584	10377		
In addition, pools of salasted cloner, as well as libraries containing appoints alongs were					

assigned an "ES" number (internal reference) and deposited with the ATCC. Table 13 below provides the ATCC Accession Nos. of the ES deposits, all of which were deposited on or before June 13, 2000. The names of the clones contained within each of these deposits are provided in the tables numbered 22 and greater (inserted before the claims).

Table 12: Pools of Clones and Libraries Deposited with ATCC on or before June 13, 2000.

Library No.	CMCC No.	ATCC Accession No.	Library No.	CMCC No.	ATCC Accession No.
ES168	5276	PTA-2027	ES 189	5304	PTA-2052
ES169	5277	PTA-2028	ES 190	5305	PTA-2053
ES170	5284	PTA-2029	ES 191	5306	PTA-2054
ES171	5285	PTA-2030	ES 192	5307	PTA-2055

Library No.	CMCC No.	ATCC Accession No.	Library No.	CMCC No.	ATCC Accession No.
ES172	5286	PTA-2031	ES 193	5308	PTA-2056
ES173	5287	PTA-2032	ES 194	5309	PTA-2057
ES174	5288	PTA-2033	ES 195	5310	PTA-2058
ES175	5289	PTA-2034	ES 196	5311	PTA-2059
ES176	5290	PTA-2035	ES 197	5312	PTA-2060
ES177	5291	PTA-2036	ES 198	5313	PTA-2061
ES178	5292	PTA-2037	ES 199	5314	PTA-2062
ES179	5293	PTA-2038	ES 200	5315	PTA-2048
ES180	5294	PTA-2039	ES 201	5316	PTA-2049
ES181	5295	PTA-2040	ES 202	5317	PTA-2063
ES182	5296	PTA-2041	ES 203	5318	PTA-2064
ES183	5297	PTA-2042	ES 204	5319	PTA-2065
ES184	5298	PTA-2043	ES 205	5320	PTA-2066
ES185	5299	PTA-2044	ES 206	5321	PTA-2067
ES 186	5301	PTA-2045	ES 207	5322	PTA-2068
ES 187	5302	PTA-2046	ES 208	5253	PTA-2050
ES 188	5303	PTA-2047	ES 209	5324	PTA-2051

Table 13 (inserted before the claims) provides the names of the clones in each of the above libraries. Retrieval of Individual Clones from Deposit of Pooled Clones. Where the ATCC deposit is composed of a pool of cDNA clones or a library of cDNA clones, the deposit was prepared by first transfecting each of the clones into separate bacterial cells. The clones in the pool or library were then deposited as a pool of equal mixtures in the composite deposit. Particular clones can be obtained from the composite deposit using methods well known in the art. For example, a bacterial cell containing a particular clone can be identified by isolating single colonies, and identifying colonies containing the specific clone through standard colony hybridization techniques, using an oligonucleotide probe or probes designed to specifically hybridize to a sequence of the clone insert (e.g., a probe based upon unmasked sequence of the encoded polynucleotide having the indicated SEQ ID NO). The probe should be designed to have a Tm of approximately 80°C (assuming 2°C for each A or T and 4°C for each G or C). Positive colonies can then be picked, grown in culture, and the recombinant clone isolated. Alternatively, probes designed in this manner can be used to PCR to isolate a nucleic acid molecule from the pooled clones according to methods well known in the art, e.g., by purifying the cDNA from the deposited culture pool, and using the probes in PCR reactions to produce an amplified product having the corresponding desired polynucleotide sequence.

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Table 2

1	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
1	498509	2102,H02.gz43_275066	F	M00063514B:E04	GRRpz
2	644927	2102.F08.gz43_275160	F	M00063493D:G07	GRRpz
3	621081	2102.G10.gz43_275193	F	M00063507B:B08	GRRpz
4	558900	2102.F14.gz43_275256	F	M00063496B;F07	GRRpz
5	464791	2102,A16.gz43_275283	F	M00063157B:B09	GRRpz
6	558147	2102.B18.gz43_275316	F	M00063457A:B12	GRRpz
7	379154	2102.I24.gz43_275419	F	M00063601D:C05	GRRpz
8	643609	2103.F01.gz43_275432	F	M00063866B:H02	GRRpz
9	377696	2103.M06.gz43_275519	F	M00063943B:G12	GRRpz
10	554395	2103,K10.gz43_275581	F	M00063927A:B08	GRRpz
11	402353	2103.B13.gz43_275620	F	M00063803B:F11	GRRpz
12	380514	2103.E17.gz43_275687	F	M00063864A:H10	GRRpz
13	147279	2103.J21.gz43_275756	F	M00063923B:A04	GRRpz
14	884	2128.K06.gz43_277507	F	M00028627B:F12	MV-522
15	188377	2128.H15.gz43_277648	F	M00028359D:F09	MV-522
16	155615	2128.P17.gz43_277688	F	M00032476D:F07	MV-522
17	125604	2128.P18.gz43_277704	F	M00032477A:B02	MV-522
18	432159	2128.A24.gz43_277785	F	M00028061D:D10	MV-522
19	429735	2130.N02.gz43_277847	F	M00032736A:B06	MV-522
20	72838	2130.005.gz43_277896	F	M00032745C:F03	MV-522
21	436888	2130.N09.gz43_277959	F	M00032738D:G11	MV-522
22	427571	2130.H13.gz43_278017	F	M00032685B:C10	MV-522
23	38494	2130.H16.gz43_278065	F	M00032686C:D10	MV-522
24	78607	2130.H18.gz43_278097	F	M00032686D:G09	MV-522
25	90192	2130,J18,gz43_278099	F	M00032703D:E10	MV-522
26	44615	2130,I20,gz43_278130	F	M00032695B:A01	MV-522
27	376753	2130,M21.gz43_278150	F	M00032732A;A03	MV-522
28	6342	2152.H10.gz43_278370	F	M00039109A:H09	UCP-3
29	387530	2152.D12.gz43_278398	F	M00039069D:E12	UCP-3
30	36453	2152.B14.gz43_278428	F	M00039052B:H03	UCP-3
31	376044	2152.A18.gz43_278491	F	M00039047C:A05	UCP-3
32	375415	2152.J18.gz43_278500	F	M00039121C:C06	UCP-3
33	375706	2153.F03.gz43_278756	F	M00039222C:G06	UCP-3
34	379046	2153.J06.gz43_278808	F	M00039244C:F10	UCP-3
35	390017	2153.I08.gz43_278839	F	M00039240A;H08	UCP-3
36	377596	2153.008.gz43_278845	F	M00039274C:D12	UCP-3
37	376384	2153.I11.gz43_278887	F	M00039241A:H11	UCP-3
38	372952	2153.K14.gz43 278937	F	M00039251A;G12	UCP-3
39	377696	2154.M04.gz43_279163	F	M00039404B:G11	UCP-3
40	60037	2154.G05.gz43_279173	F	M00039330B:H09	UCP-3
41	390968	2154.I18.gz43_279383	F	M00039346C:G08	UCP-3
42	376044	2154.G21.gz43 279429	F	M00039336A;C07	UCP-3

Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	Т	CLONE ID	LIBRARY
43	60037	2155.J03.gz43 279528	F	M00039632A:C01	UCP-3
44	376839	2155.N05.gz43 279564	F	M00039664B:H10	UCP-3
45	373680	2155.P11.gz43 279662	F	M00039674B:G11	UCP-3
46	373973	2155.C12.gz43 279665	F	M00039457A:D07	UCP-3
47	379805	2155.N15.gz43 279724	F	M00039665C:B01	UCP-3
48	376439	2155.F16.gz43_279732	F	M00039479A:A10	UCP-3
49	376994	2155.I16.gz43_279735	F	M00039619C:B01	UCP-3
50	378420	2155.B21.gz43_279808	F	M00039451B:F01	UCP-3
51	375510	2155.F21.gz43_279812	F	M00039479D:C06	UCP-3
52	24210	2155.P22.gz43_279838	F	M00039675C:C05	UCP-3
53	397167	2164.J03.gz43_279912	F	M00039778D:D05	UCP-3
54	380025	2164.I10.gz43_280023	F	M00039774A:E11	UCP-3
55	216179	2164.P10.gz43_280030	F	M00039823D:D09	UCP-3
56	379046	2164.B15.gz43_280096	F	M00039681B:F05	UCP-3
57	379279	2164.A24.gz43_280239	F	M00039679C:A02	UCP-3
58	376074	2164.024.gz43_280253	F	M00039820D:F08	UCP-3
59	373905	2165.H01.gz43_280262	F	M00039883D:G06	UCP-3
60	1300	2165.K01.gz43_280265	F	M00039905B:F09	UCP-3
61	393635	2165.H06.gz43_280342	F	M00039885B:A10	UCP-3
62	396969	2165.P07.gz43_280366	F	M00039978D:C04	UCP-3
63	398061	2165.F09.gz43_280388	F	M00039871C;C01	UCP-3
64	376808	2165.I10.gz43_280407	F	M00039893A:G12	UCP-3
65	6342	2166.A06.gz43_281279	F	M00039986B:A11	UCP-3
66	233814	2166.K07.gz43_281305	F	M00040076B;D01	UCP-3
67	24210	2166.N07.gz43_281308	F	M00040096D:C03	UCP-3
68	24210	2166,A11.gz43_281359	F	M00039988B:C08	UCP-3
69	377696	2166.J11.gz43_281368	F	M00040070C:D11	UCP-3
70	398831	2166.A23.gz43_281551	F	M00040001A;H02	UCP-3
71	185432	2166.P24.gz43_281582	F	M00040113C:H09	UCP-3
72	505275	2104.D01.gz43_297323	F	M00064082C:D11	GRRpz
73	11379	2104.N01.gz43_297333	F	M00064194B:A02	GRRpz
74	446397	2104.N03.gz43_297365	F	M00064195C;B02	GRRpz
75	418763	2104.P04.gz43_297383	F	M00064220B;E01	GRRpz
76	469367	2104.N05.gz43_297397	F	M00064196D:C10	GRRpz
77	639578	2104.C08.gz43_297434	F	M00064077B:H02	GRRpz
78	376559	2104.E08.gz43 297436	F	M00064101B;E12	GRRpz
79	649035	2104.D11.gz43_297483	F	M00064085B;A12	GRRpz
80	647586	2104.H11.gz43_297487	F	M00064131B:A09	GRRpz
81	646187	2104.F12.gz43_297501	F	M00064112A:G03	GRRpz
82	727888	2104.N13.gz43_297525	F	M00064201A:C08	GRRpz
83	558382	2104.E17.gz43_297580	F	M00064104B;A01	GRRpz
84	52644	2104.C19.gz43_297610	F	M00064081A:D04	GRRpz

Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	Т	CLONE ID	LIBRARY
85	454622	2104.I20.gz43 297632	F	M00064147B:G08	GRRpz
86	554032	2104.A21.gz43 297640	F	M00064064D:D11	GRRpz
87	463217	2104.E23.gz43 297676	F	M00064107C:E03	GRRpz
88	189073	1521.H06.gz43 303496	F	M00027103D:B05	MCF-7
89	188309	1521.J06.gz43 303498	F	M00027219B:G12	MCF-7
90	187525	1521.N10.gz43_303566	F	M00027514C:F01	MCF-7
91	215366	1521.N12.gz43_303598	F	M00027517C:F08	MCF-7
92	186594	1521.H13.gz43_303608	F	M00027123D:F02	MCF-7
93	218904	1521.O13.gz43_303615	F	M00027586B:B03	MCF-7
94	189993	1521.C14.gz43_303619	F	M00023406A:G03	MCF-7
95	222818	1521.H14.gz43 303624	F	M00027126C:H05	MCF-7
96	185056	1521.O16.gz43_303663	F	M00027587C:F02	MCF-7
97	186404	1521.P20.gz43_303728	F	M00027694C:C11	MCF-7
98	649744	2116.I06.gz43_306217	F	M00063580A:A07	WOca
99	535955	2116.P08.gz43_306256	F	M00063994B:D10	WOca
100	375328	2116.J09.gz43_306266	F	M00063592B:E09	WOca
101	449206	2116.A13.gz43_306321	F	M00063165C:F10	WOca
102	89082	2116.E15.gz43_306357	F	M00063552B:B06	WOca
103	730600	2116.D21.gz43_306452	F	M00063546B:F01	WOca
104	372621	2116.O22.gz43_306479	F	M00063988A:C02	WOca
105	416886	2117.M03.gz43_306557	F	M00064394A:C02	WOca
106	372621	2117.D05.gz43_306580	F	M00064307C:E12	WOca
107	644919	2118.N03.gz43_306942	F	M00064601C:H06	WOca
108	446397	2118.A09.gz43_307025	F	M00064446D:C08	WOca
109	177443	2118.M09.gz43_307037	F	M00064592D:F05	WOca
110	730238	2118.H24.gz43_307272	F	M00064534D:H04	WOca
111	427907	2131.A01.gz43_307885	F	M00032766A:A10	MV-522
112	48238	2131.M02.gz43_307913	F	M00032886A:D04	MV-522
113	226324	2131.B04.gz43_307934	F	M00032783A:H08	MV-522
114	441801	2131.E06.gz43_307969	F	M00032809B:E10	MV-522
115	62016	2131.K11.gz43_308055	F	M00032872B:A02	MV-522
116	48238	2131.E12.gz43_308065	F	M00032811A:G10	MV-522
117	34071	2131.I13.gz43_308085	F	M00032857A:B02	MV-522
118	221686	2131.B14.gz43_308094	F	M00032786A:H04	MV-522
119	440284	2131.J16.gz43_308134	F	M00032865A;D11	MV-522
120	12481	2131.P18.gz43_308172	F	M00032914B:D09	MV-522
121	20453	2131.A19.gz43_308173	F	M00032779A:A04	MV-522
122	26926	2131.I19.gz43_308181	F	M00032858D:H11	MV-522
123	37805	2131.P19.gz43_308188	F	M00032915B:D01	MV-522
124	441874	2131.I21.gz43_308213	F	M00032859C:E04	MV-522
125	48238	2131.A23.gz43_308237	F	M00032780A:B09	MV-522
126	169458	1513.O03.gz43_300292	F	M00022648C:D08	MDA-MB-231

Table 2

	able 2				
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ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
127	145815	1513.F09.gz43 300379	F	M00022135D:D06	MDA-MB-231
128	158321	1513.H09.gz43 300381	F	M00022255B:F12	MDA-MB-231
129	168195	1513.L10.gz43_300401	F	M00022537D:C05	MDA-MB-231
130	101499	1513.J13.gz43 300447	F	M00022445D:E12	MDA-MB-231
131	142842	1513.A14.gz43_300454	F	M00007960D:E09	MDA-MB-231
132	153316	1513.E14.gz43_300458	F	M00022072A:E12	MDA-MB-231
133	142614	1513.A15.gz43_300470	F	M00007963D:D03	MDA-MB-231
134	99011	1513.N16.gz43_300499	F	M00022622A:G01	MDA-MB-231
135	171073	1513.P18.gz43_300533	F	M00022710C:H03	MDA-MB-231
136	120049	1513.D19.gz43_300537	F	M00021925A:H07	MDA-MB-231
137	446572	1562.B01.gz43_207804	F	M00042546A;D03	UC2-NormColon
138	464091	1562.F01.gz43_207808	F	M00042552D:A11	UC2-NormColon
139	408386	1562.J01.gz43_207812	F	M00042560B:A01	UC2-NormColon
140	446829	1562.L02.gz43_207830	F	M00042563A;F10	UC2-NormColon
141	43338	1562.P03.gz43_207850	F	M00042569B:G07	UC2-NormColon
142	451780	1562.F04.gz43_207856	F	M00042554A:C02	UC2-NormColon
143	456462	1562.J04.gz43 207860	F	M00042560B:G10	UC2-NormColon
144	469511	1562.G05.gz43 207873	F	M00042555D:G10	UC2-NormColon
145	455075	1562.H05.gz43 207874	F	M00042557A:D09	UC2-NormColon
146	447346	1562.P05.gz43 207882	F	M00042569C:B05	UC2-NormColon
147	147196	1562.M06.gz43 207895	F	M00042564D:F10	UC2-NormColon
148	467262	1562.F08.gz43_207920	F	M00042554C:E02	UC2-NormColon
149	459536	1562.L09.gz43_207942	F	M00042563C:D08	UC2-NormColon
150	460190	1562.P09.gz43_207946	F	M00042569D:D02	UC2-NormColon
151	447597	1562.F10.gz43_207952	F	M00042554C:F09	UC2-NormColon
152	451737	1562.G10.gz43_207953	F	M00042556A:G12	UC2-NormColon
153	446614	1562.L10.gz43_207958	F	M00042563C:E02	UC2-NormColon
154	432159	1562.M10.gz43_207959	F	M00042565A:G05	UC2-NormColon
155	459523	1562.L11.gz43_207974	F	M00042563D:D02	UC2-NormColon
156	460516	1562.P11.gz43_207978	F	M00042570A:E08	UC2-NormColon
157	446674	1562.D12.gz43_207982	F	M00042550A:D12	UC2-NormColon
158	465594	1562.F12.gz43_207984	F	M00042554D:C08	UC2-NormColon
159	466719	1562.G12.gz43_207985	F	M00042556B:D12	UC2-NormColon
160	463487	1562.M12.gz43_207991	F	M00042565A:H03	UC2-NormColon
161	446389	1562.B13.gz43_207996	F	M00042546D:E06	UC2-NormColon
162	462149	1562.C14.gz43_208013	F	M00042548B:G01	UC2-NormColon
163	447174	1562.G14.gz43_208017	F	M00042556B:E10	UC2-NormColon
164	457405	1562.K14.gz43_208021	F	M00042562C:A07	UC2-NormColon
165	460766	1562.O15.gz43_208041	F	M00042568C:E08	UC2-NormColon
166	446703	1562.N16.gz43_208056	F	M00042567C:E02	UC2-NormColon
167	446981	1562.D18.gz43_208078	F	M00042550C:H10	UC2-NormColon
168	387077	1562.H18.gz43 208082	F	M00042558A:D03	UC2-NormColon

Table 2

	able 2				
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169	460972	1562.N18.gz43_208088	F	M00042567C:E07	UC2-NormColon
170	417078	1562.P18.gz43_208090	F	M00042570B:F11	UC2-NormColon
171	461559	1562.L19.gz43 208102	F	M00042564A:F01	UC2-NormColon
172	464905	1562.F20.gz43 208112	F	M00042555B:B07	UC2-NormColon
173	468330	1562.H20.gz43 208114	F	M00042558A:F11	UC2-NormColon
174	446620	1562,J21.gz43_208132	F	M00042561B:E12	UC2-NormColon
175	457842	1562.P21.gz43_208138	F	M00042570C:B12	UC2-NormColon
176	446531	1562.B22.gz43_208140	F	M00042547B:D11	UC2-NormColon
177	446621	1562.J24.gz43_208180	F	M00042561C:E12	UC2-NormColon
178	380127	1562.N24.gz43_208184	F	M00042567D:E12	UC2-NormColon
179	464275	1563.D01.gz43_208190	F	M00042575D:A11	UC2-NormColon
180	456920	1563.C04.gz43_208237	F	M00042574C:A04	UC2-NormColon
181	466971	1563.B05.gz43_208252	F	M00042573A:D05	UC2-NormColon
182	142559	1563.J05.gz43_208260	F	M00042698A:A01	UC2-NormColon
183	457890	1563.G08.gz43_208305	F	M00042691D:B03	UC2-NormColon
184	453605	1563.K08.gz43_208309	F	M00042699C:H06	UC2-NormColon
185	465339	1563.N08.gz43_208312	F	M00042705A:B07	UC2-NormColon
186	461734	1563.I09.gz43_208323	F	M00042695C:F05	UC2-NormColon
187	459961	1563.D12.gz43_208366	F	M00042576B:D11	UC2-NormColon
188	463951	1563.F13.gz43_208384	F	M00042580C:A03	UC2-NormColon
189	468783	1563.A14.gz43_208395	F	M00042571C:F03	UC2-NormColon
190	447597	1563.E14.gz43_208399	F	M00042578A:E08	UC2-NormColon
191	463368	1563.I14.gz43_208403	F	M00042695D:H01	UC2-NormColon
192	451383	1563.C15.gz43_208413	F	M00042575A:E02	UC2-NormColon
193	467293	1563.G16.gz43_208433	F	M00042692A:E11	UC2-NormColon
194	446922	1563.O16.gz43_208441	F	M00042707C:A09	UC2-NormColon
195	446213	1563.D17.gz43_208446	F	M00042576D:A08	UC2-NormColon
196	460244	1563.F19.gz43_208480	F	M00042691A:D08	UC2-NormColon
197	460789	1563.A20.gz43_208491	F	M00042572B:E05	UC2-NormColon
198	451382	1563.C20.gz43_208493	F	M00042575B:F02	UC2-NormColon
199	468109	1563.K20.gz43_208501	F	M00042700C:F11	UC2-NormColon
200	447326	1563.L20.gz43_208502	F	M00042702B:B07	UC2-NormColon
201	463896	1563.B21.gz43_208508	F	M00042573D:A10	UC2-NormColon
202	446839	1563.D21.gz43_208510	F	M00042576D:F01	UC2-NormColon
203	446933	1563.E22.gz43_208527	F	M00042579A:B05	UC2-NormColon
204	447826	1563.L22.gz43_208534	F	M00042702B:G07	UC2-NormColon
205	446409	1563.B23.gz43_208540	F	M00042574A:F05	UC2-NormColon
206	453908	1573.N01.gz43_208584	F	M00042895C:C10	UC2-NormColon
207	447421	1573.D02,gz43_208590	F	M00042747D:C08	UC2-NormColon
208	447141	1573.P02.gz43_208602	F	M00042898A:H05	UC2-NormColon
209	447645	1573.E04.gz43_208623	F	M00042750D:E07	UC2-NormColon
210	401426	1573.K06.gz43_208661	F	M00042890D:G05	UC2-NormColon

Table 2

SEQ ID NO CLUSTER SEQ NAME T CLONE ID LIBRARY	T	able 2				
211		OV TYGONIA	GT0 1/11			
212		CHUBILL				LIBRARY
213 633946 1573.M08.gz43 208695 F M00042894B:E05 UC2-NormColon 214 557974 1573.L09.gz43 208718 F M00042889C:E03 UC2-NormColon 215 466902 1573.D10.gz43 208718 F M00042788D:D08 UC2-NormColon 216 642146 1573.F10.gz43 208720 F M00042788D:D08 UC2-NormColon 217 464205 1573.F10.gz43 208724 F M000427889D:A12 UC2-NormColon 218 641890 1573.L10.gz43 208726 F M00042889D:A12 UC2-NormColon 219 650195 1573.H11.gz43 208738 F M00042886C:F01 UC2-NormColon 220 467293 1573.F12.gz43 208738 F M00042886C:F01 UC2-NormColon 221 650756 1573.H11.gz43 208736 F M00042886C:F01 UC2-NormColon 222 452611 1573.N13.gz43 208766 F M00042886C:F03 UC2-NormColon 222 452611 1573.N13.gz43 208766 F M00042896A:E03 UC2-NormColon 222 452611 1573.N13.gz43 208796 F M00042886C:F01 UC2-NormColon 222 452611 1573.N14.gz43 208799 F M00042886C:F01 UC2-NormColon 222 452612 1573.N14.gz43 208800 F M00042886C:F03 UC2-NormColon 222 452612 1573.H14.gz43 208818 F M00042886D:E10 UC2-NormColon 225 645690 1573.F15.gz43 208800 F M00042886D:E10 UC2-NormColon 226 645470 1573.H16.gz43 208818 F M00042886D:E10 UC2-NormColon 227 651029 1573.M17.gz43 208839 F M000428810-G50 UC2-NormColon 228 46238 1573.K19.gz43 208889 F M000428810-G50 UC2-NormColon 229 486238 1573.K19.gz43 208896 F M00042891C-H01 UC2-NormColon 232 472101 1574.O11.gz43 208896 F M00042891C-H01 UC2-NormColon 233 641875 1574.P02.gz43 208996 F M00042891C-H01 UC2-NormColon 234 465104 1574.003.gz43 208996 F M000429916-G1 UC2-NormColon 236 470641 1574.003.gz43 208996 F M000429916-G1 UC2-NormColon 237 479641 1574.003.gz43 208996 F M00042908-B10 UC2-NormColon 238 470462 1574.B06.gz43 209006 F M00042908-B10 UC2-NormColon 240 641525 1574.B06.gz43 209006 F M00042908-B10 UC2-NormColon 240 641625 1574.B06.						
214 557974 1573.L199_gz43_208710 F M00042892C:E03 UC2-NormColon 1574.D10_gz43_208718 F M00042748D:D08 UC2-NormColon 1574.D10_gz43_208720 F M00042882C:F06 UC2-NormColon 1574.D10_gz43_208726 F M00042885C:F06 UC2-NormColon 1574.D10_gz43_208726 F M00042885C:F06 UC2-NormColon 1574.D10_gz43_208726 F M00042885C:F06 UC2-NormColon 1574.D10_gz43_208738 F M00042885C:F06 UC2-NormColon 1574.D10_gz43_208738 F M00042885C:F01 UC2-NormColon 1574.D10_gz43_208738 F M00042885C:F01 UC2-NormColon 1574.D10_gz43_208736 F M00042885C:F01 UC2-NormColon 1574.D10_gz43_208736 F M00042885C:F01 UC2-NormColon 1574.D10_gz43_208796 F M00042885C:F01 UC2-NormColon 1574.D10_gz43_208796 F M00042885C:F01 UC2-NormColon 1574.D10_gz43_208796 F M00042885C:F03 UC2-NormColon 1574.D10_gz43_208830 F M00042885D:D10 UC2-NormColon 1574.D10_gz43_208830 F M00042885D:D10 UC2-NormColon 1574.D10_gz43_208830 F M00042885D:D10 UC2-NormColon 1573.D11_gz43_208830 F M00042885D:D10 UC2-NormColon 1573.D11_gz43_208899 F M00042885D:D10 UC2-NormColon 1573.D11_gz43_208899 F M00042885D:D10 UC2-NormColon 1574.D11_gz43_208899 F M00042885D:D10 UC2-NormColon 1574.D11_gz43_208999 F M00042891D:D10 UC2-NormColon 1574.D11_gz43_208999 F M00042891D:D10 UC2-NormColon 1574.D11_gz43_208999 F M00042891D:D10 UC2-NormColon 1574.D11_gz43_208999 F M00042891D:D10 UC2-NormColon 1574.D11_gz43_208999 F M0004290B:B10 UC2-NormC						
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217						
218					M00042882C:F06	UC2-NormColon
219 650195 1573.H11.gz43_208738 F M00042886C:F01 UC2-NormColon 220 467293 1573.F12.gz43_208752 F M00042884C:G07 UC2-NormColon 221 650756 1573.N12.gz43_208760 F M00042886C:F01 UC2-NormColon 222 452611 1573.N13.gz43_208760 F M00042896A:E03 UC2-NormColon 223 650944 1573.N14.gz43_208790 F M00042896A:E03 UC2-NormColon 224 639372 1573.E15.gz43_208799 F M00042886C:C11 UC2-NormColon 225 645690 1573.F15.gz43_208890 F M00042881C:C11 UC2-NormColon 225 645690 1573.F15.gz43_208819 F M00042881C:C11 UC2-NormColon 227 651029 1573.M17.gz43_208839 F M00042881C:G10 UC2-NormColon 227 651029 1573.M17.gz43_208839 F M00042881C:G10 UC2-NormColon 228 639849 1573.F18.gz43_208848 F M00042881A:F06 UC2-NormColon 229 486238 1573.K19.gz43_208899 F M00042881A:F06 UC2-NormColon 230 463060 1573.G12.gz43_208897 F M00042881A:G09 UC2-NormColon 231 469150 1573.C2.gz43_208999 F M00042876.G09 UC2-NormColon 232 472101 1574.O01.gz43_208996 F M00042747A:G12 UC2-NormColon 233 641875 1574.P02.gz43_208996 F M00042747A:G12 UC2-NormColon 234 465104 1574.A03.gz43_208986 F M00042791C:G11 UC2-NormColon 235 470641 1574.A03.gz43_208986 F M0004290B:B10 UC2-NormColon 236 451624 1574.D04.gz43_209006 F M0004290B:B10 UC2-NormColon 237 639391 1574.N04.gz43_209006 F M0004290B:B10 UC2-NormColon 238 470462 1574.B04.gz43_209006 F M0004290B:B10 UC2-NormColon 240 641525 1574.B05.gz43_209028 F M0004290B:B00 UC2-NormColon 241 642691 1574.B06.gz43_209002 F M00042915A:E06 UC2-NormColon 242 466697 1574.G06.gz43_209004 F M00042915A:G01 UC2-NormColon 243 649965 1574.B06.gz43_209004 F M00042915A:G01 UC2-NormColon 244 465065 1574.B06.gz43_209004 F M00042915A:G01 UC2-NormColon 244 466697 1574.G06.gz43_209004 F M00042915A:G01 UC2-NormColon 244 466697 1574.G06.gz						
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232 472101 1574.001 gz43 208969 F M00054797D:F01 UC2-NormColon 233 641875 1574.P02 gz43 208986 F M00054799C:G11 UC2-NormColon 234 465104 1574.003 gz43 208987 F M00042909B:B10 UC2-NormColon 235 470641 1574.D04 gz43 208906 F M00042909C:A10 UC2-NormColon 236 451624 1574.D04 gz43 209016 F M00042906C:A10 UC2-NormColon 237 639391 1574.N04 gz43 209020 F M00042908B:H01 UC2-NormColon 239 482043 1574.N05 gz43 209020 F M00042908B:H01 UC2-NormColon 240 641525 1574.N05 gz43 209039 F M00042908A:B01 UC2-NormColon 241 642691 1574.E06 gz43 209093 F M00042910A:B01 UC2-NormColon 242 466697 1574.B06 gz43 209044 F M0004291DA:C0 UC2-NormColon 243 649965 1574.M06 gz43 209044 F M0004291CA:C01 UC2-NormColon	230	463060		F_	M00042885A:G09	UC2-NormColon
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236	234	465104	1574.A03.gz43_208987	F	M00042900B:B10	UC2-NormColon
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239 482043 1574_J05_gz43_209028 F M00042915A:E06 UC2-NormColon	237	639391	1574.N04.gz43_209016	F	M00054796B:A01	UC2-NormColon
240 641525 1574.N05.gz43_209032 F M00054796B:C08 UC2-NormColon 241 642691 1574.E06.gz43_209039 F M00042910D:A02 UC2-NormColon 242 466697 1574.E06.gz43_209041 F M00042910D:A02 UC2-NormColon 243 649965 1574.H06.gz43_209042 F M00042910A:C01 UC2-NormColon 244 651051 1574.J06.gz43_209047 F M00042915A:G10 UC2-NormColon 245 450506 1574.M06.gz43_209047 F M00054794C:G11 UC2-NormColon 246 433572 1574.D06.gz43_209049 F M00042902C:E11 UC2-NormColon 247 447660 1574.B7.2gz43_209053 F M00042902C:E11 UC2-NormColon 248 447147 1574.C07.gz43_209053 F M00042909B:C04 UC2-NormColon 249 647639 1574.E08.gz43_209056 F M00042909B:C04 UC2-NormColon 250 649965 1574.G08.gz43_209091 F M000429143:B05 UC2-NormColon <td< td=""><td>238</td><td>470462</td><td>1574.B05.gz43_209020</td><td>F</td><td>M00042902B:H01</td><td>UC2-NormColon</td></td<>	238	470462	1574.B05.gz43_209020	F	M00042902B:H01	UC2-NormColon
241 642691 1574.E06.gz43 209039 F M00042908A:B01 UC2-NormColon 242 466697 1574.G06.gz43 209041 F M00042910D:A02 UC2-NormColon 243 649965 1574.H06.gz43 209042 F M00042912A:C01 UC2-NormColon 244 651051 1574.J06.gz43 209044 F M00042915A:G10 UC2-NormColon 245 450506 1574.M06.gz43 209047 F M00034798E:A01 UC2-NormColon 246 453572 1574.O06.gz43 209042 F M00042902:B11 UC2-NormColon 247 447660 1574.B07.gz43 209052 F M00042902:B11 UC2-NormColon 248 447147 1574.C07.gz43 209053 F M00042905A:A07 UC2-NormColon 249 647639 1574.F07.gz43 209053 F M00042909B:C04 UC2-NormColon 250 649965 1574.G08.gz43 209073 F M00042910D:E11 UC2-NormColon 251 639371 1574.109.gz43 209091 F M00042910D:E1 UC2-NormColon	239	482043	1574.J05.gz43_209028	F	M00042915A:E06	UC2-NormColon
242 466697 1574.G06.gzd3_209041 F M00042910D:A02 UC2-NormColon 243 649965 1574.H06.gzd3_209042 F M00042912A:C01 UC2-NormColon 244 651051 1574.H06.gzd3_209044 F M00042915A:G10 UC2-NormColon 245 450506 1574.M06.gzd3_209047 F M00054794C:G11 UC2-NormColon 246 453572 1574.006.gzd3_209049 F M00042916A:B01 UC2-NormColon 247 447660 1574.B07.gzd3_209052 F M00042905A:A07 UC2-NormColon 248 447147 1574.C07.gzd3_209053 F M00042905A:A07 UC2-NormColon 249 647639 1574.F07.gzd3_209056 F M00042909B:C04 UC2-NormColon 250 649965 1574.G08.gzd3_209073 F M00042910D:E11 UC2-NormColon 251 639371 1574.109.gzd3_209091 F M00042910A:B05 UC2-NormColon	240	641525	1574.N05.gz43_209032	F	M00054796B:C08	UC2-NormColon
243 649965 1574.H06 gz43 209042 F M00042912A:C01 UC2-NormColon 244 651051 1574.J06 gz43 209044 F M00042915A:G10 UC2-NormColon 245 450506 1574.M06 gz43 209047 F M00043794:G11 UC2-NormColon 246 453572 1574.006 gz43 209099 F M00043794:G11 UC2-NormColon 247 447660 1574.B07 gz43 209053 F M00042905A:A07 UC2-NormColon 248 447147 1574.C07 gz43 209053 F M00042905A:A07 UC2-NormColon 249 647639 1574.F07 gz43 209056 F M00042909B:C04 UC2-NormColon 250 649965 1574.G08.gz43 209073 F M00042914:B05 UC2-NormColon 251 639371 1574.109 gz43 209091 F M00042914:B05 UC2-NormColon	241	642691	1574.E06.gz43_209039	F	M00042908A:B01	UC2-NormColon
244 651051 1574J06gg43 209044 F M00042915A:G10 UC2-NormColon 245 450506 1574.M06.gz43 209047 F M00034794C:G11 UC2-NormColon 246 453572 1574.006.gz43 209049 F M00054798B:A01 UC2-NormColon 247 447660 1574.B07.gz43 209052 F M00042902C:B11 UC2-NormColon 248 447147 1574.C07.gz43 209053 F M00042909B:C04 UC2-NormColon 249 647639 1574.F07.gz43 209056 F M00042909B:C04 UC2-NormColon 250 649965 1574.G08.gz43 209073 F M00042910D:E11 UC2-NormColon 251 639371 1574.109.gz43 209091 F M00042910A:B05 UC2-NormColon	242	466697	1574.G06.gz43_209041	F	M00042910D:A02	UC2-NormColon
245 450506 1574.M06.gz43_209047 F M00054794C:G11 UC2-NormColon 246 453372 1574.O06.gz43_209049 F M00054798B:A01 UC2-NormColon 247 447660 1574.B07.gz43_209052 F M00042902E11 UC2-NormColon 248 447147 1574.C07.gz43_209053 F M00042905A:A07 UC2-NormColon 249 647639 1574.F07.gz43_209056 F M00042909B:C04 UC2-NormColon 250 649965 1574.G08.gz43_209073 F M00042910D:E11 UC2-NormColon 251 639371 1574.109.gz43_209091 F M000429143:B05 UC2-NormColon	243	649965	1574.H06.gz43 209042	F	M00042912A:C01	UC2-NormColon
246 453572 1574.006.gzd3_209049 F M00054798B:A01 UC2-NormColon 247 447660 1574.B07.gzd3_209052 F M00042902C:E11 UC2-NormColon 248 447147 1574.C07.gzd3_209053 F M000429005A:A07 UC2-NormColon 249 647639 1574.F07.gzd3_209056 F M00042909B:C04 UC2-NormColon 250 649965 1574.G08.gzd3_209073 F M00042910D:E11 UC2-NormColon 251 639371 1574.109.gzd3_209091 F M00042914A:B05 UC2-NormColon	244	651051	1574.J06.gz43 209044	F	M00042915A:G10	UC2-NormColon
247 447660 1574.B07.gz43_209052 F M00042902C:E11 UC2-NormColon 248 447147 1574.C07.gz43_209053 F M000429095A:A07 UC2-NormColon 249 647639 1574.F07.gz43_209056 F M00042909B:C04 UC2-NormColon 250 649965 1574.G08.gz43_209073 F M00042910D:E11 UC2-NormColon 251 639371 1574.109.gz43_209091 F M00042914A:B05 UC2-NormColon	245	450506	1574.M06.gz43 209047	F	M00054794C:G11	UC2-NormColon
248 447147 1574.C07.gz43 209053 F M00042905A:A07 UC2-NormColon 249 647639 1574.F07.gz43 209056 F M00042909B:C04 UC2-NormColon 250 649965 1574.G08.gz43 209073 F M00042910D:E11 UC2-NormColon 251 639371 1574.109.gz43 209091 F M00042914A:B05 UC2-NormColon	246	453572	1574.O06.gz43 209049	F	M00054798B:A01	UC2-NormColon
249 647639 1574.F07.g243 209056 F M00042909B:C04 UC2-NormColon 250 649965 1574.G08.gz43 209073 F M00042910D:E11 UC2-NormColon 251 639371 1574.I09.gz43 209091 F M00042914A:B05 UC2-NormColon	247		1574.B07.gz43_209052	F	M00042902C:E11	UC2-NormColon
249 647639 1574.F07.gz43_209056 F M00042909B:C04 UC2-NormColon 250 649965 1574.G08.gz43_209073 F M00042910D:E11 UC2-NormColon 251 639371 1574.I09.gz43_209091 F M00042914A:B05 UC2-NormColon	248	447147	1574.C07.gz43_209053	F	M00042905A:A07	UC2-NormColon
250 649965 1574.G08.gz43_209073 F M00042910D:E11 UC2-NormColon 251 639371 1574.I09.gz43_209091 F M00042914A:B05 UC2-NormColon	249	647639		F		
251 639371 1574.I09.gz43_209091 F M00042914A:B05 UC2-NormColon	250	649965				
	252	474298		F	M00054794D:D02	UC2-NormColon

Table 2

1	able 2				
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	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
253	646568	1574,M10,gz43 209111	F	M00054794D:D08	UC2-NormColon
254	639787	1574.P10.gz43 209114	F	M00054800B;C06	UC2-NormColon
255	447750	1574.A11.gz43 209115	F	M00042901A:F12	UC2-NormColon
256	649810	1574.G11.gz43 209121	F	M00042911A;A02	UC2-NormColon
257	645924	1574.H11.gz43 209122	F	M00042912B:F11	UC2-NormColon
258	413767	1574.L11.gz43 209126	F	M00054793C:D11	UC2-NormColon
259	641069	1574.E12.gz43_209135	F	M00042908B:A11	UC2-NormColon
260	649900	1574.G12.gz43 209137	F	M00042911A:B02	UC2-NormColon
261	560868	1574.J12.gz43 209140	F	M00042915C:E05	UC2-NormColon
262	513262	1574.P12.gz43 209146	F	M00054800B:C11	UC2-NormColon
263	640306	1574.L13.gz43 209158	F	M00054793D:H11	UC2-NormColon
264	389591	1574.A14.gz43_209163	F	M00042901A:H11	UC2-NormColon
265	447815	1574.C14.gz43 209165	F	M00042905C:G08	UC2-NormColon
266	95617	1574.D14.gz43 209166	F	M00042907A:B11	UC2-NormColon
267	465984	1574.B15.gz43_209180	F	M00042903B:C09	UC2-NormColon
268	447692	1574.D15.gz43_209182	F	M00042907A:F03	UC2-NormColon
269	641029	1574.G15.gz43_209185	F	M00042911A:D04	UC2-NormColon
270	650527	1574.J15.gz43_209188	F	M00042915D:A11	UC2-NormColon
271	641315	1574.O15.gz43_209193	F	M00054798D:A12	UC2-NormColon
272	474298	1574.P15.gz43_209194	F	M00054800B:E08	UC2-NormColon
273	645954	1574.H16.gz43_209202	F	M00042912D:H08	UC2-NormColon
274	640181	1574.M16.gz43_209207	F	M00054795A:A08	UC2-NormColon
275	650235	1574.F17.gz43_209216	F	M00042909C:F10	UC2-NormColon
276	640356	1574.H17.gz43_209218	F	M00042912D:H10	UC2-NormColon
277	649744	1574.E18.gz43_209231	F	M00042908C:A03	UC2-NormColon
278	650564	1574.F18.gz43_209232	F	M00042909D:B11	UC2-NormColon
279	527355	1574.G18.gz43_209233	F	M00042911A:H12	UC2-NormColon
280	452989	1574.H18.gz43_209234	F	M00042913A:D09	UC2-NormColon
281	648159	1574.K18.gz43_209237	F	M00054792B:A03	UC2-NormColon
282	714629	1574.E20.gz43_209263	F	M00042908C:D12	UC2-NormColon
283	467364	1574.I20.gz43_209267	F	M00042914B:H03	UC2-NormColon
284	644789	1574.N20.gz43_209272	F	M00054797C:F03	UC2-NormColon
285	646404	1574.J21.gz43_209284	F	M00054791A:G04	UC2-NormColon
286	639740	1574.H22.gz43_209298	F	M00042913B:E10	UC2-NormColon
287	649852	1574.I22.gz43_209299	F	M00042914D:B10	UC2-NormColon
288	644376	1574.G23.gz43_209313	F	M00042911B:H08	UC2-NormColon
289	639901	1574.P23.gz43_209322	F	M00054800D:D08	UC2-NormColon
290	647086	1574.E24.gz43_209327	F	M00042908D:G12	UC2-NormColon
291	562247	1574.G24.gz43_209329	F	M00042911C:D01	UC2-NormColon
292	446974	1574.J24.gz43_209332	F	M00054791B:C09	UC2-NormColon
293	513248	1575.A01.gz43_209339	F	M00054800D:F08	UC2-NormColon
294	467989	1575.M01.gz43_209351	F	M00055429B;B12	UC2-NormColon

Table 2

	able 2				
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ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
295	650184	1575.N01.gz43_209352	F	M00055430D:F04	UC2-NormColon
296	649872	1575.E02.gz43_209359	F	M00054916A:A05	UC2-NormColon
297	649506	1575.G02.gz43_209361	F	M00054918A:D02	UC2-NormColon
298	446254	1575.M02.gz43 209367	F	M00055429B:E12	UC2-NormColon
2 99	641542	1575.O02.gz43_209369	F	M00055432B:H02	UC2-NormColon
300	649933	1575.I03.gz43_209379	F	M00054920B:C04	UC2-NormColon
301	639444	1575.M03.gz43_209383	F	M00055429B:G04	UC2-NormColon
302	553100	1575.G04.gz43_209393	F	M00054918A:F09	UC2-NormColon
303	549699	1575.A05.gz43_209403	F	M00054911A:C08	UC2-NormColon
304	639256	1575.E05.gz43_209407	F	M00054916A:E05	UC2-NormColon
305	644556	1575.H05.gz43_209410	F	M00054919C:F06	UC2-NormColon
306	500337	1575.I05.gz43_209411	F	M00054920B:C12	UC2-NormColon
307	643924	1575.K05.gz43_209413	F	M00055427A:F01	UC2-NormColon
308	639394	1575.E06.gz43_209423	F	M00054916A:F10	UC2-NormColon
309	452986	1575.J06.gz43_209428	F	M00055426B:B02	UC2-NormColon
310	654723	1575.K06.gz43_209429	F	M00055427A:F02	UC2-NormColon
311	645344	1575.N07.gz43_209448	F	M00055431A:H05	UC2-NormColon
312	645162	1575.O07.gz43_209449	F	M00055432C:F01	UC2-NormColon
313	589098	1575.B08.gz43_209452	F	M00054912B:C05	UC2-NormColon
314	643843	1575.F08.gz43_209456	F	M00054917C:D03	UC2-NormColon
315	647109	1575.P08.gz43_209466	F	M00055434A:A03	UC2-NormColon
316	467381	1575.D09.gz43_209470	F	M00054914D:G07	UC2-NormColon
317	570812	1575.N09.gz43_209480	F	M00055431B:A01	UC2-NormColon
318	559776	1575.A10.gz43_209483	F	M00054911A:G01	UC2-NormColon
319	648532	1575.F10.gz43_209488	F	M00054917C:F03	UC2-NormColon
320	449861	1575.G10.gz43_209489	F	·M00054918B:H01	UC2-NormColon
321_	643843	1575.I10.gz43_209491	F	M00054920C:A06	UC2-NormColon
322_	41141	1575.K10.gz43_209493	F	M00055427B:E01	UC2-NormColon
323	648664	1575.E11.gz43_209503	F	M00054916B:E02	UC2-NormColon
324	640814	1575,K11,gz43_209509	F	M00055427B:F06	UC2-NormColon
325	584071	1575.M11.gz43_209511	F	M00055429D:G07	UC2-NormColon
326	450225	1575.C12.gz43_209517	F	M00054913C:G03	UC2-NormColon
327	452707	1575.I12.gz43_209523	F	M00054920C:D05	UC2-NormColon
328	243722	1575.L12.gz43_209526	F	M00055428C:G06	UC2-NormColon
329	641057	1575.B13.gz43_209532 -	F	M00054912C:C01	UC2-NormColon
330	413767	1575.D13.gz43_209534	F	M00054915A:G03	UC2-NormColon
331	504568	1575.K13.gz43_209541	F	M00055427C:A06	UC2-NormColon
332	639992	1575.A15.gz43_209563	F	M00054911B:E10	UC2-NormColon
333	456923	1575.G15.gz43_209569	F	M00054918D:C03	UC2-NormColon
334	649555	1575.I15.gz43_209571	F	M00054920C:F02	UC2-NormColon
335	649746	1575.J15.gz43_209572	F	M00055426C:C10	UC2-NormColon
336	644692	1575.F16.gz43_209584	F	M00054917D:A12	UC2-NormColon

Table 2

	able 2				
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ID NO	CLUSTER	SEQ NAME	Т	CLONE ID	LIBRARY
337	452204	1575.G16.gz43_209585	F	M00054918D:C11	UC2-NormColon
338	61616	1575.H16.gz43_209586	F	M00054920A:A05	UC2-NormColon
339	495143	1575.M16.gz43 209591	F	M00055430B:E08	UC2-NormColon
340	644371	1575.O16.gz43 209593	F	M00055433A:B07	UC2-NormColon
341	630269	1575.K17.gz43_209605	F	M00055427C:E12	UC2-NormColon
342	446932	1575.O17.gz43_209609	F	M00055433A:B08	UC2-NormColon
343	639662	1575.F18.gz43_209616	F	M00054917D:D12	UC2-NormColon
344	611927	1575.L18.gz43_209622	F	M00055428D:G12	UC2-NormColon
345	641925	1575.O18.gz43_209625	F	M00055433A:C02	UC2-NormColon
346	651020	1575.P18.gz43_209626	F	M00055434C:B11	UC2-NormColon
347	639255	1575.F19.gz43_209632	F	M00054917D:E05	UC2-NormColon
348	468222	1575.G19.gz43_209633	F	M00054918D:H09	UC2-NormColon
349	468783	1575.K19.gz43_209637	F	M00055427C:H11	UC2-NormColon
350	417130	1575.B20.gz43_209644	F	M00054912D:G01	UC2-NormColon
351	448606	1575.B21.gz43_209660	F	M00054912D:G04	UC2-NormColon
352	456420	1575.K21.gz43_209669	F	M00055427D:E05	UC2-NormColon
353	640369	1575.P21.gz43_209674	F	M00055434D:B06	UC2-NormColon
354	473854	1575,E22.gz43_209679	F	M00054917A:F07	UC2-NormColon
355	564440	1575.H22.gz43_209682	F	M00054920A:C11	UC2-NormColon
356	639520	1575.L22.gz43_209686	F	M00055429A:H04	UC2-NormColon
357	649195	1575.O22.gz43_209689	F	M00055433A:E04	UC2-NormColon
358	57183	1575.P22.gz43_209690	F	M00055434D:E09	UC2-NormColon
359	648609	1575.E23.gz43_209695	F	M00054917B:A05	UC2-NormColon
360	648532	1575.F23.gz43_209696	F	M00054917D:H02	UC2-NormColon
361	642073	1575.N23.gz43_209704	F	M00055432B:B04	UC2-NormColon
362	377855	1575.B24.gz43_209708	F	M00054913A:B12	UC2-NormColon
363	452845	1575.G24.gz43_209713	F	M00054919A:H04	UC2-NormColon
364	638758	1575.O24.gz43_209721	F	M00055433C:A11	UC2-NormColon
365	639829	1576.A04.gz43_209771	F	M00055435B:C09	UC2-NormColon
366	484145	1576.I05.gz43_209795	F	M00055446B:A12	UC2-NormColon
367	640055	1576.A08.gz43_209835	F	M00055435C:E12	UC2-NormColon
368	639928	1576,I08.gz43_209843	F	M00055446B:D08	UC2-NormColon
369	526606	1576.L08.gz43_209846	F	M00055450B:G07	UC2-NormColon
370	653616	1576.E09.gz43_209855	F	M00055440D:D02	UC2-NormColon
371	624440	1576.M12.gz43_209911	F	M00055451C:B08	UC2-NormColon
372	560791	1576.O12.gz43_209913	F	M00055453D:E12	UC2-NormColon
373	639287	1576.A13.gz43_209915	F	M00055435D:G11	UC2-NormColon
374	475184	1576.M14.gz43_209943	F	M00055451C:E10	UC2-NormColon
375	626061	1576.A15.gz43_209947	F	M00055436A:E04	UC2-NormColon
376	640734	1576.E16.gz43_209967	F	M00055441A:G07	UC2-NormColon
377	466092	1576.B17.gz43_209980	F	M00055437D:B06	UC2-NormColon
378	639711	1576.A19.gz43 210011	F	M00055436B:B06	UC2-NormColon

Table 2

	able 2				
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ID NO	CLUSTER	SEQ NAME	Т	CLONE ID	LIBRARY
379	640522	1576,C21.gz43 210045	F	M00055439B:B07	UC2-NormColon
380	651038	1585,I01.gz43 210211	F	M00055465A:C05	UC2-NormColon
381	466440	1585.B02.gz43_210220	F	M00055456C;A10	UC2-NormColon
382	648123	1585.C02.gz43_210221	F	M00055458B;B04	UC2-NormColon
383	646318	1585.K02.gz43_210229	F	M00055468A:A05	UC2-NormColon
384	598589	1585,E03.gz43_210239	F	M00055460B:G06	UC2-NormColon
385	447863	1585.F03.gz43_210240	F	M00055461A:H03	UC2-NormColon
386	562236	1585.P05.gz43_210282	F	M00055494C;G10	UC2-NormColon
387	559369	1585,A06.gz43_210283	F	M00055456A:B03	UC2-NormColon
388	484126	1585.B06.gz43_210284	F	M00055456C:G04	UC2-NormColon
389	558839	1585.P06.gz43_210298	F	M00055494C:G11	UC2-NormColon
390	555820	1585.J07.gz43_210308	F	M00055467A:A07	UC2-NormColon
391	544461	1585.H09.gz43_210338	F	M00055463D:G01	UC2-NormColon
392	642478	1585.A10.gz43_210347	F	M00055456A:F01	UC2-NormColon
393	643843	1585.C11.gz43_210365	F	M00055458C:G02	UC2-NormColon
394	640400	1585.G12.gz43_210385	F	M00055462C:A11	UC2-NormColon
395	645201	1585.L12.gz43_210390	F	M00055469C:F09	UC2-NormColon
396	640792	1585.F13.gz43_210400	F	M00055461C:E05	UC2-NormColon
397	640913	1585.H13.gz43_210402	F	M00055464B:E06	UC2-NormColon
398	639607	1585,I13.gz43_210403	F	M00055465D:F12	UC2-NormColon
399	639932	1585.B14.gz43_210412	F	M00055456D:G04	UC2-NormColon
400	640662	1585.D14.gz43_210414	F	M00055460A:D01	UC2-NormColon
401	598589	1585.G14.gz43_210417	F	M00055462C:C03	UC2-NormColon
402	556654	1585.H14.gz43_210418	F	M00055464B;E11	UC2-NormColon
403	555193	1585.O14.gz43_210425	F	M00055493D:B07	UC2-NormColon
404	641066	1585.E15.gz43_210431	F	M00055460D:B06	UC2-NormColon
405	664711	1585.H15.gz43_210434	F	M00055464B:G03	UC2-NormColon
406	549611	1585.I15.gz43_210435	F	M00055466A:C05	UC2-NormColon
407	639726	1585.K16.gz43_210453	F	M00055468C:B07	UC2-NormColon
408	607422	1585.F17.gz43_210464	F	M00055461D:C09	UC2-NormColon
409	584745	1585.M17.gz43_210471	F	M00055491A;H01	UC2-NormColon
410	661194	1585.H19.gz43_210498	F	M00055464D:A04	UC2-NormColon
411	559549	1585.J20.gz43_210516	F	M00055467D:A01	UC2-NormColon
412	641645	1585.A22.gz43_210539	F	M00055456B:G08	UC2-NormColon
413	641467	1585.C22.gz43_210541	F	M00055459B:A02	UC2-NormColon
414	640368	1585.F22.gz43_210544	F	M00055462A:A09	UC2-NormColon
415	412416	1585.G22.gz43_210545	F	M00055462D:H12	UC2-NormColon
416	650914	1585.H22.gz43 210546	F	M00055464D.F08	UC2-NormColon
417	3	1585.J22.gz43_210548	F	M00055467D:C10	UC2-NormColon
418	646318	1585.N22.gz43_210552	F	M00055492C:H07	UC2-NormColon
419	237288	1585.P22.gz43_210554	F	M00055495C:F03	UC2-NormColon
420	650605	1585.B23.gz43_210556	F	M00055457D:F09	UC2-NormColon

Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
421	643991	1585.E23.gz43_210559	F	M00055461A;A06	UC2-NormColon
422	398061	1585.G23.gz43_210561	F	M00055463A:A11	UC2-NormColon
423	505933	1585.J24.gz43_210580	F	M00055467D:G08	UC2-NormColon
424	640068	1585.K24.gz43_210581	F	M00055469A:D08	UC2-NormColon
425	474821	1585.N24.gz43_210584	F	M00055492D:B08	UC2-NormColon
426	234606	1587.C01.gz43_211361	F	M00055519A:F08	UC2-NormColon
427	597780	1587.E01.gz43_211363	F	M00055521C:B08	UC2-NormColon
428	492483	1587.B02.gz43_211376	F	M00055517C:H07	UC2-NormColon
429	644051	1587.C03.gz43_211393	F	M00055519A:H01	UC2-NormColon
430	639056	1587,E03.gz43_211395	F	M00055521C:C08	UC2-NormColon
431	643723	1587.L03.gz43_211402	F	M00055529D:D05	UC2-NormColon
432	639459	1587.N03.gz43_211404	F	M00055532C:G08	UC2-NormColon
433	639826	1587.P03.gz43_211406	F	M00055535B:A11	UC2-NormColon
434	552783	1587.A04.gz43_211407	F	M00055516B:E11	UC2-NormColon
435	590218	1587.D04.gz43_211410	F	M00055520B:D11	UC2-NormColon
436	559324	1587.E04.gz43_211411	F	M00055521C:D02	UC2-NormColon
437	646711	1587.L04.gz43_211418	F	M00055529D:D11	UC2-NormColon
438	639132	1587.M04.gz43_211419	F	M00055531B:D10	UC2-NormColon
439	207552	1587.N04.gz43_211420	F	M00055532D:A12	UC2-NormColon
440	116869	1587.004.gz43_211421	F	M00055533D:G02	UC2-NormColon
441	481220	1587.D05.gz43_211426	F	M00055520B:E04	UC2-NormColon
442	645544	1587.K05.gz43_211433	F	M00055528D:B02	UC2-NormColon
443	650617	1587,M05.gz43_211435	F	M00055531B:E05	UC2-NormColon
444	194095	1587.005.gz43_211437	F	M00055534A:E06	UC2-NormColon
445	446984	1587.B06.gz43_211440	F	M00055517D:D09	UC2-NormColon
446	290226	1587.B07.gz43_211456	F	M00055517D:D11	UC2-NormColon
447	634409	1587.P07.gz43_211470	F	M00055535C:A03	UC2-NormColon
448	640419	1587.G08.gz43_211477	F	M00055524B:B08	UC2-NormColon
449	649149	1587.L09.gz43_211498	F	M00055530A:C07	UC2-NormColon
450	640072	1587.P10.gz43_211518	F	M00055535C:E08	UC2-NormColon
451	641615	1587.J11.gz43_211528	F	M00055528A:E08	UC2-NormColon
452	639213	1587.M13.gz43_211563	F	M00055531D:E06	UC2-NormColon
453	641216	1587.N15.gz43_211596	F	M00055533B:B11	UC2-NormColon
454	650161	1587.A16.gz43_211599	F	M00055517A:D09	UC2-NormColon
455	558785	1587.B23.gz43_211712	F	M00055519A:C01	UC2-NormColon
456	642631	1588.B02.gz43_211780	F	M00055538C:E04	UC2-NormColon
457	639543	1588.F02.gz43_211784	F	M00055543D:H03	UC2-NormColon
458	473701	1588.F03.gz43_211800	F	M00055544A:A07	UC2-NormColon
459	562274	1588.G03.gz43_211801	F	M00055545B:A04	UC2-NormColon
460	643277	1588.D05.gz43_211830	F	M00055541C:D02	UC2-NormColon
461	556867	1588.G05.gz43_211833	F	M00055545B:C01	UC2-NormColon
462	646350	1588.P05.gz43_211842	F	M00055556A:A04	UC2-NormColon

Table 2

1 1					
SEO			ORIEN		
l	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
463	649335	1588.B06.gz43 211844	F	M00055538D:D12	UC2-NormColon
464	646159	1588.J06.gz43 211852	F	M00055548B:C03	UC2-NormColon
465	649099	1588.P06.gz43 211858	F	M00055556A:E07	UC2-NormColon
466	32812	1588.H09.gz43 211898	F	M00055546C:F11	UC2-NormColon
467	639480	1588.J09.gz43 211900	F	M00055548B:G06	UC2-NormColon
468	644242	1588.N09.gz43_211904	F	M00055553B:H04	UC2-NormColon
469	644723	1588.N11.gz43 211936	F	M00055553C:D06	UC2-NormColon
470	648748	1588.F13.gz43 211960	F	M00055544C;A01	UC2-NormColon
471	639099	1588,G13.gz43 211961	F	M00055545C:D09	UC2-NormColon
472	562414	1588.F15.gz43 211992	F	M00055544C:B07	UC2-NormColon
473	641078	1588.N15.gz43 212000	F	M00055553C:H12	UC2-NormColon
474	505042	1588.G16.gz43 212009	F	M00055545D:B06	UC2-NormColon
475	640747	1588.N16.gz43 212016	F	M00055553D:C07	UC2-NormColon
476	453606	1588,D17.gz43 212022	F	M00055542B:B11	UC2-NormColon
477	419706	1588.A18.gz43 212035	F	M00055538A:C05	UC2-NormColon
478	557983	1588.H21.gz43 212090	F	M00055547B:C05	UC2-NormColon
479	639114	1588.D22.gz43 212102	F	M00055542C:D07	UC2-NormColon
480	442531	1588,J22,gz43 212108	F	M00055549A:B04	UC2-NormColon
481	649668	1597.N02.gz43 212176	F	M00055574D:E02	UC2-NormColon
482	639781	1597.D05.gz43 212214	F	M00055561C:C07	UC2-NormColon
483	639056	1597.E05.gz43 212215	F	M00055562D:B06	UC2-NormColon
484	639698	1597,P05.gz43 212226	F	M00055577C:G02	UC2-NormColon
485	469731	1597.B06.gz43 212228	F	M00055559A:D06	UC2-NormColon
486	639593	1597.G06.gz43_212233	F	M00055566A:E02	UC2-NormColon
487	557975	1597.J06.gz43 212236	F	M00055569D:C02	UC2-NormColon
488	640356	1597.E08.gz43 212263	F	M00055563A:A02	UC2-NormColon
489	647431	1597.E09.gz43 212279	F	M00055563A:C03	UC2-NormColon
490	31112	1597.C10.gz43_212293	F	M00055560D:C03	UC2-NormColon
491	454825	1597.F10.gz43_212296	F	M00055565A:C08	UC2-NormColon
492	555021	1597.B12.gz43 212324	F	M00055559C:D07	UC2-NormColon
493	650740	1597.I12.gz43 212331	F	M00055569A:D01	UC2-NormColon
494	659031	1597.J12.gz43 212332	F	M00055570B:F07	UC2-NormColon
495	592122	1597.D14.gz43_212358	F	M00055562B:D03	UC2-NormColon
496	45921	1597.N14.gz43 212368	F	M00055575C:B04	UC2-NormColon
497	556511	1597,H15,gz43 212378	F	M00055568A:B04	UC2-NormColon
498	637966	1597.A16.gz43 212387	F	M00055558B:G11	UC2-NormColon
499	553318	1597.F16.gz43_212392	F	M00055565A:F11	UC2-NormColon
500	23961	1597.C17.gz43_212405	F	M00055561A:F07	UC2-NormColon
501	650235	1597.M17.gz43_212415	F	M00055574B:F06	UC2-NormColon
502	596882	1597.F18.gz43_212424	F	M00055565B:F04	UC2-NormColon
503	644928	1597.I18.gz43_212427	F	M00055569A:F06	UC2-NormColon
504	589483	1597.L18.gz43_212430	F	M00055573B:D11	UC2-NormColon

Table 2

	able 2				
SEO			ORIEN	1	
ID NO	CLUSTER	SEQ NAME	Т	CLONE ID	LIBRARY
505	640956	1597.N18.gz43 212432	F	M00055575D:G04	UC2-NormColon
506	553602	1597.L19.gz43 212446	F	M00055573B:E01	UC2-NormColon
507	640851	1597.O20.gz43 212465	F	M00055577A:G09	UC2-NormColon
508	460666	1597.A21.gz43 212467	F	M00055558D:C08	UC2-NormColon
509	640195	1597.D22.gz43 212486	F	M00055562C:F02	UC2-NormColon
510	668852	1597.P22.gz43 212498	F	M00055578C:F11	UC2-NormColon
511	645913	1597.A23.gz43_212499	F	M00055558D:D07	UC2-NormColon
512	446230	1597.G23.gz43 212505	F	M00055567A:A06	UC2-NormColon
513	649349	1597.N23.gz43_212512	F	M00055576A:F07	UC2-NormColon
514	645538	1597.C24.gz43 212517	F	M00055561B:G12	UC2-NormColon
515	647069	1597.G24.gz43 212521	F	M00055567A:A11	UC2-NormColon
516	638869	1597.H24.gz43_212522	F	M00055568C:F07	UC2-NormColon
517	556	1597.I24.gz43_212523	F	M00055569B:G10	UC2-NormColon
518	648483	1597.024.gz43_212529	F	M00055577B:F05	UC2-NormColon
519	447272	1598,M01.gz43_212543	F	M00055595C:F12	UC2-NormColon
520	640025	1598.B02.gz43 212548	F	M00055581A:C02	UC2-NormColon
521	415538	1598.E02.gz43_212551	F	M00055585B:F01	UC2-NormColon
522	478229	1598.M02.gz43_212559	F	M00055595C:G03	UC2-NormColon
523	644572	1598.H03.gz43_212570	F	M00055589B:E08	UC2-NormColon
524	648580	1598.B04.gz43 212580	F	M00055581C:A01	UC2-NormColon
525	470769	1598.L04.gz43_212590	F	M00055594C:B03	UC2-NormColon
526	554273	1598.M04.gz43_212591	F	M00055595D:C04	UC2-NormColon
527	641262	1598.C05.gz43_212597	F	M00055583A:A05	UC2-NormColon
528	645707	1598.F05.gz43_212600	F	M00055586D:G07	UC2-NormColon
529	463028	1598.N05.gz43_212608	F	M00055597B:B04	UC2-NormColon
530	640695	1598,P05.gz43_212610	F	M00055601B:D12	UC2-NormColon
531	641925	1598.D06.gz43_212614	F	M00055584A:G11	UC2-NormColon
532	449247	1598.E06.gz43_212615	F	M00055585C:F05	UC2-NormColon
533	646590	1598,L06,gz43 212622	F	M00055594C:F11	UC2-NormColon
534	641191	1598.A07.gz43_212627	F	M00055579C:D04	UC2-NormColon
535	641191	1598.P07.gz43_212642	F	M00055601B:H02	UC2-NormColon
536	641838	1598.K09.gz43_212669	F	M00055593A:F08	UC2-NormColon
537	641440	1598.N09.gz43_212672	F	M00055597D:B05	UC2-NormColon
538	640826	1598.M10.gz43_212687	F	M00055596A:E07	UC2-NormColon
539	459521	1598.A11.gz43_212691	F	M00055579D:C11	UC2-NormColon
540	140909	1598.G11.gz43_212697	F	M00055588B:H11	UC2-NormColon
541	641467	1598.J12.gz43_212716	F	M00055592B:C10	UC2-NormColon
542	455864	1598.L12.gz43_212718	F	M00055594D:F07	UC2-NormColon
543	641957	1598.A13.gz43_212723	F	M00055579D:G09	UC2-NormColon
544	638941	1598.K14.gz43_212749	F	M00055593C:D08	UC2-NormColon
545	648159	1598.B15.gz43_212756	F	M00055582B:A06	UC2-NormColon
546	509973	1598.G15.gz43_212761	F	M00055588C:G09	UC2-NormColon

Table 2

T	able 2				
SEQ			ORIEN		
	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
547	641425	1598.A17.gz43 212787	F	M00055580B;B08	UC2-NormColon
548	647312	1598.O17.gz43 212801	F	M00055600C;C02	UC2-NormColon
549	86311	1598.C18.gz43_212805	F	M00055583C:A01	UC2-NormColon
550	515931	1598.O19.gz43 212833	F	M00055600D:B02	UC2-NormColon
551	651000	1598.D21.gz43 212854	F	M00055585A;E12	UC2-NormColon
552	553087	1598.E21.gz43 212855	F	M00055586C;A06	UC2-NormColon
553	646914	1598.G21.gz43 212857	F	M00055589A:B06	UC2-NormColon
554	446371	1598.K21.gz43_212861	F	M00055594B:A01	UC2-NormColon
555	640298	1598.N23.gz43 212896	F	M00055599D:C08	UC2-NormColon
556	201904	1599.A01.gz43 212923	F	M00055602B:B10	UC2-NormColon
557	641338	1599.B01.gz43 212924	F	M00055603D:A09	UC2-NormColon
558	640221	1599.L01.gz43 212934	F	M00055619C:F07	UC2-NormColon
559	454527	1599.F02.gz43 212944	F	M00055610B:E04	UC2-NormColon
560	640868	1599.I08.gz43 213043	F	M00055615C:E01	UC2-NormColon
561	641069	1599.N09.gz43 213064	F	M00055623D:G05	UC2-NormColon
562	559380	1599.B10.gz43 213068	F	M00055604D:F05	UC2-NormColon
563	641680	1599.D10.gz43 213070	F	M00055608C:E03	UC2-NormColon
564	639703	1599.I11.gz43 213091	F	M00055615D:C07	UC2-NormColon
565	641542	1599.C12.gz43 213101	F	M00055606D:C05	UC2-NormColon
566	5201	1599.A15.gz43 213147	F	M00055602D:G08	UC2-NormColon
567	607715	1599.D15.gz43 213150	F	M00055608C:G11	UC2-NormColon
568	640416	1599.L17.gz43 213190	F	M00055620D;D05	UC2-NormColon
569	554833	1599.B19.gz43 213212	F	M00055606A;B11	UC2-NormColon
570	639480	1599.K19.gz43 213221	F	M00055619B:H04	UC2-NormColon
571	607138	1599.P21.gz43 213258	F	M00055628B;B07	UC2-NormColon
572	561626	1599 L23.gz43 213286	F	M00055621B:G03	UC2-NormColon
573	548959	1600.I01.gz43 213315	F	M00055639D:D03	UC2-NormColon
574	639934	1600.H02.gz43_213330	F	M00055638D:D07	UC2-NormColon
575	640956	1600.I03.gz43 213347	F	M00055639D:F08	UC2-NormColon
576	554722	1600.L03.gz43 213350	F	M00055644A:D12	UC2-NormColon
577	649717	1600.D04.gz43 213358	F	M00055633D:A02	UC2-NormColon
578	646695	1600,H04,gz43 213362	F	M00055638D:E09	UC2-NormColon
579	639886	1600,N04,gz43 213368	F	M00055647C:D02	UC2-NormColon
580	451615	1600.K05.gz43 213381	F	M00055643A:C01	UC2-NormColon
581	635965	1600.I07.gz43 213411	F	M00055640A:G03	UC2-NormColon
582	379040	1600,L07.gz43 213414	F	M00055644B:H12	UC2-NormColon
583	640799	1600.C10.gz43 213453	F	M00055632D:A06	UC2-NormColon
584	642246	1600.K11.gz43 213477	F	M00055643B:E05	UC2-NormColon
585	467563	1600.N11.gz43 213480	F	M00055647D:B11	UC2-NormColon
586	640320	1600.011.gz43 213481	F	M00055649A:H07	UC2-NormColon
587	557401	1600.I12.gz43 213491	F	M00055640C:E06	UC2-NormColon
588	646352	1600.M13.gz43 213511	F	M00055646C:B04	UC2-NormColon

Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
589	645004	1600.B15.gz43_213532	F	M00055630B:G04	UC2-NormColon
590	641702	1600.P17.gz43_213578	F	M00055651A:E06	UC2-NormColon
591	562000	1600.I19.gz43_213603	F	M00055641A:C12	UC2-NormColon
592	453441	1600.N19.gz43_213608	F	M00055648B:C01	UC2-NormColon
593	641210	1600,M20.gz43_213623	F	M00055647B:A05	UC2-NormColon
594	458736	1600.P20.gz43_213626	F	M00055651B:F08	UC2-NormColon
595	553285	1600.O21.gz43_213641	F	M00055650A:B05	UC2-NormColon
596	455075	1600.A22.gz43_213643	F	M00055629B:G09	UC2-NormColon
597	166041	1600.D23.gz43_213662	F	M00055634C;F09	UC2-NormColon
598	553516	1600.J23.gz43_213668	F	M00055642D:A05	UC2-NormColon
599	559423	1600.B24.gz43_213676	· F	M00055632A:B11	UC2-NormColon
600	131348	1600.C24.gz43_213677	F	M00055633B:G02	UC2-NormColon
601	650180	1600.E24.gz43_213679	F	M00055636A:H12	UC2-NormColon
602	642361	1600.F24.gz43_213680	F	M00055637B:A01	UC2-NormColon
603	561069	1600.K24.gz43_213685	F	M00055643D:G11	UC2-NormColon
604	550515	1600.P24.gz43_213690	F	M00055651C:E01	UC2-NormColon
605	559854	1693.D01.gz43_213694	F	M00055819A:B10	UC2-NormColon
606	650579	1693.J01.gz43_213700	F	M00055827A:A12	UC2-NormColon
607	648567	1693.L02.gz43_213718	F	M00055829C:A07	UC2-NormColon
608	645155	1693.H03.gz43_213730	F	M00055825B:C11	UC2-NormColon
609	650204	1693.H04.gz43_213746	F	M00055825B:E03	UC2-NormColon
610	643054	1693.H06.gz43_213778	F	M00055825B:F09	UC2-NormColon
611	640464	1693.I06.gz43_213779	F	M00055826A:G04	UC2-NormColon
612	363172	1693.A07.gz43_213787	F	M00055816D:A10	UC2-NormColon
613	674526	1693.G08.gz43_213809	F	M00055823B:F02	UC2-NormColon
614	481864	1693.N08.gz43_213816	F	M00055832A:A08	UC2-NormColon
615	492242	1693.A09.gz43_213819	F	M00055816D:B11	UC2-NormColon
616	638837	1693.C10.gz43_213837	F	M00055818B:H03	UC2-NormColon
617	517280	1693.K10.gz43_213845	F	M00055828B:E10	UC2-NormColon
618	452863	1693.A13.gz43_213883	F	M00055816D:E10	UC2-NormColon
619	642962	1693.H13.gz43_213890	F	M00055825D:A03	UC2-NormColon
620	645746	1693.I13.gz43_213891	F	M00055826B:G07	UC2-NormColon
621	645146	1693.L15.gz43_213926	F	M00055829D:H10	UC2-NormColon
622	644781	1693.M15.gz43_213927	F	M00055831A:C06	UC2-NormColon
623	643397	1693.A17.gz43_213947	F	M00055817A:E05	UC2-NormColon
624	645073	1693.L17.gz43_213958	F	M00055830A:G10	UC2-NormColon
625	648580	1693.A19.gz43_213979	F	M00055817A:H07	UC2-NormColon
626	650487	1693.I19.gz43_213987	F	M00055826C:G06	UC2-NormColon
627	455716	1693.J19.gz43_213988	F	M00055827D:A01	UC2-NormColon
628	551681	1693.H20.gz43_214002	F	M00055825D:D11	UC2-NormColon
629	642054	1693.P21.gz43_214026	F	M00055835C:F08	UC2-NormColon
630	556286	1693,J22.gz43_214036	F	M00055827D:C02	UC2-NormColon

Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
631	449542	1693.A23.gz43_214043	F	M00055817B:C04	UC2-NormColon
632	517237	1693.M23.gz43_214055	F	M00055831B:C04	UC2-NormColon
633	647191	1694.J01.gz43_214084	F	M00055846B:F11	UC2-NormColon
634	644212	1694.G04.gz43_214129	F	M00055843B;D10	UC2-NormColon
635	549624	1694.P04.gz43_214138	F	M00055854C:C07	UC2-NormColon
636	645149	1694.A05.gz43_214139	F	M00055836A:B12	UC2-NormColon
637	981	1694.F05.gz43_214144	F	M00055842B:A04	UC2-NormColon
638	647448	1694.I05.gz43_214147	F	M00055845C:A11	UC2-NormColon
639	570939	1694.P05.gz43_214154	F	M00055854C:E03	UC2-NormColon
640	547841	1694.L06.gz43_214166	F	M00055849C:G07	UC2-NormColon
641	466265	1694.M06.gz43_214167	F	M00055851A:C03	UC2-NormColon
642	649842	1694.M07.gz43_214183	F	M00055851A:C09	UC2-NormColon
643	468222	1694.I08.gz43_214195	F	M00055845C:C12	UC2-NormColon
644	450949	1694.L08.gz43_214198	F	M00055849D:B04	UC2-NormColon
645	603388	1694.O08.gz43_214201	F	M00055853C:C12	UC2-NormColon
646	485237	1694.E09.gz43_214207	F	M00055841B:F09	UC2-NormColon
647	644063	1694.F09.gz43_214208	F	M00055842B;D04	UC2-NormColon
648	639341	1694.N09.gz43_214216	F	M00055852A:C12	UC2-NormColon
649	471364	1694.P09.gz43_214218	F	M00055854C:H11	UC2-NormColon
650	452735	1694.A10.gz43_214219	F	M00055836C:D01	UC2-NormColon
651	503546	1694.E10.gz43_214223	F	M00055841B:H03	UC2-NormColon
652	646420	1694.I10.gz43_214227	F	M00055845C:E02	UC2-NormColon
653	647577	1694.O10.gz43_214233	F	M00055853C;H03	UC2-NormColon
654	504944	1694.B11.gz43_214236	F	M00055837D:G10	UC2-NormColon
655	522869	1694.D11.gz43_214238	F	M00055840B:B02	UC2-NormColon
656	644548	1694.F11.gz43_214240	F	M00055842C:A11	UC2-NormColon
657	644314	1694.H11.gz43_214242	F	M00055844D:E12	UC2-NormColon
658	650492	1694.O11.gz43_214249	F	M00055853D;A07	UC2-NormColon
659	643800	1694.B12.gz43_214252	F	M00055838A:A03	UC2-NormColon
660	643843	1694.E12.gz43_214255	F	M00055841C:A03	UC2-NormColon
661	448450	1694.B13.gz43_214268	F	M00055838A:B02	UC2-NormColon
662	643804	1694.C13.gz43_214269	F	M00055839B;A10	UC2-NormColon
663	469511	1694.L13.gz43_214278	F	M00055849D:H09	UC2-NormColon
664	643130	1694.O13.gz43_214281	F	M00055853D:B04	UC2-NormColon
665	640171	1694.J14.gz43_214292	F	M00055846D:G11	UC2-NormColon
666	647522	1694.M14.gz43_214295	F	M00055851A:H10	UC2-NormColon
667	642293	1694.N14.gz43_214296	F	M00055852B:G09	UC2-NormColon
668	404816	1694.P14.gz43_214298	F	M00055855A:B11	UC2-NormColon
669	447597	1694.A15.gz43_214299	F	M00055837A:B08	UC2-NormColon
670	467901	1694.B15.gz43_214300	F	M00055838B:D06	UC2-NormColon
671	648039	1694.C15.gz43_214301	F	M00055839B:C07	UC2-NormColon
672	607430	1694.E15.gz43_214303	F	M00055841C:D05	UC2-NormColon

Table 2

1	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	Т	CLONE ID	LIBRARY
673	645505	1694.F15.gz43_214304	F	M00055842D:C02	UC2-NormColon
674	472119	1694.G15.gz43_214305	F	M00055843D:H01	UC2-NormColon
675	645197	1694.K15.gz43_214309	F	M00055848C:A02	UC2-NormColon
676	650217	1694,O15.gz43_214313	F	M00055853D:C07	UC2-NormColon
677	644210	1694.A16.gz43_214315	F	M00055837A:D09	UC2-NormColon
678	466697	1694.H16.gz43_214322	F	M00055844D:H09	UC2-NormColon
679	650276	1694.I16.gz43_214323	F	M00055845D:G11	UC2-NormColon
680	562229	1694.M16.gz43_214327	F	M00055851B:B09	UC2-NormColon
681	462659	1694.P16.gz43_214330	F	M00055855A:G05	UC2-NormColon
682	644494	1694.B17.gz43_214332	F	M00055838B:H04	UC2-NormColon
683	24730	1694.D17.gz43_214334	F	M00055840C:D06	UC2-NormColon
684	599759	1694.E17.gz43_214335	F	M00055841C:D11	UC2-NormColon
685	15001	1694.F17.gz43_214336	F	M00055842D:D07	UC2-NormColon
686	558981	1694.K17.gz43_214341	F	M00055848C:G07	UC2-NormColon
687	592122	1694.O17.gz43_214345	F	M00055854A:B07	UC2-NormColon
688	489249	1694.A18.gz43_214347	F	M00055837A:F02	UC2-NormColon
689	447002	1694.B18.gz43_214348	F	M00055838C:A08	UC2-NormColon
690	558890	1694.C18.gz43_214349	F	M00055839B:E07	UC2-NormColon
691	644548	1694.D18.gz43_214350	F	M00055840C:H06	UC2-NormColon
692	643825	1694.G18.gz43_214353	F	M00055844A:D03	UC2-NormColon
693	645289	1694.A19.gz43_214363	F	M00055837A:H08	UC2-NormColon
694	643948	1694.B19.gz43_214364	F	M00055838C:B08	UC2-NormColon
695	537586	1694.C19.gz43_214365	F	M00055839B:H09	UC2-NormColon
696	462557	1694.F19.gz43_214368	F	M00055842D:F07	UC2-NormColon
697	647577	1694.G19.gz43_214369	F	M00055844A:D07	UC2-NormColon
698	643999	1694.H19.gz43_214370	F	M00055845A:C03	UC2-NormColon
699	651131	1694.J19.gz43_214372	F	M00055847B:G12	UC2-NormColon
700	425923	1694.M19.gz43_214375	F	M00055851B:G10	UC2-NormColon
701	639510	1694.O19.gz43_214377	F	M00055854A:D01	UC2-NormColon
702	643977	1694.D20.gz43_214382	F	M00055840D:B03	UC2-NormColon
703	140648	1694.E20.gz43_214383	F	M00055841C:H04	UC2-NormColon
704	649068	1694.G20.gz43_214385	F	M00055844A:F11	UC2-NormColon
705	644708	1694.N20.gz43_214392	. F	M00055852D:B11	UC2-NormColon
706	643909	1694.P20.gz43_214394	F	M00055855B;B11	UC2-NormColon
707	650528	1694.E21.gz43 214399	F	M00055841D:C11	UC2-NormColon
708	644569	1694,H21.gz43_214402	F	M00055845A:H10	UC2-NormColon
709	646459	1694.J21.gz43_214404	F	M00055847C:A11	UC2-NormColon
710	608873	1694,L21.gz43_214406	F	M00055850C:D01	UC2-NormColon
711	649611	1694.N21.gz43_214408	F	M00055852D:G12	UC2-NormColon
712	465576	1694.P21.gz43_214410	F	M00055855B:D12	UC2-NormColon
713	467521	1694.C22.gz43_214413	F	M00055839C:B11	UC2-NormColon
714	646810	1694.J22.gz43_214420	F	M00055847C:C01	UC2-NormColon

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Т	able 2				
SEO			ODVENI		,
ID NO	CLUSTER	CEONANG	ORIEN T	CLONETD	I IDD 4 DXI
715	647318	SEQ NAME		CLONE ID	LIBRARY
716	638854	1694.L22.gz43_214422 1694.O22.gz43_214425	F	M00055850C:G05 M00055854A:E04	UC2-NormColon
717	644987	1694.A23.gz43_214423	F	M00055837B:E07	UC2-NormColon UC2-NormColon
718	645948	1694.C23.gz43 214429	F	M00055839C:D06	UC2-NormColon
719	419751	1694,I23,gz43 214435	F	M00055846B:B12	UC2-NormColon
720	643594	1694.M23.gz43 214439	F	M00055851C:F12	UC2-NormColon
721	603388	1694.P23.gz43 214442	F	M00055855C:F11	UC2-NormColon
722	644075	1694.G24.gz43 214449	F	M00055844B:C12	UC2-NormColon
723	542282	1694.I24.gz43 214451	F	M00055846B:C12	UC2-NormColon
724	452976	1694.J24.gz43 214452	F	M00055847C:H09	UC2-NormColon
725	472801	1694.M24.gz43 214455	F	M00055851C:H05	UC2-NormColon
726	646633	1695.P03.gz43 214910	F	M00055875B:E09	UC2-NormColon
727	509027	1695.A04.gz43 214911	F	M00055855D:D12	UC2-NormColon
728	596882	1695.D05.gz43_214930	F	M00055860D:E04	UC2-NormColon
729	644442	1695.A07.gz43_214959	F	M00055855D:G08	UC2-NormColon
730	644047	1695.F07.gz43_214964	F	M00055863D:D09	UC2-NormColon
731	397399	1695.H07.gz43_214966	F	M00055866A:G10	UC2-NormColon
732	564854	1695.A08.gz43_214975	F	M00055856A:C06	UC2-NormColon
733	646372	1695.B08.gz43_214976	F	M00055857C:D09	UC2-NormColon
734	645848	1695.D08.gz43_214978	F	M00055861B:F04	UC2-NormColon
735	447035	1695.M08.gz43_214987	F	M00055872D:D12	UC2-NormColon
736	644149	1695.A09.gz43_214991	F	M00055856A:D12	UC2-NormColon
737	640147	1695.E09.gz43_214995	F	M00055862D:B02	UC2-NormColon
738	524261	1695.K09.gz43_215001	F	M00055871A:H06	UC2-NormColon
739	449500	1695.E11.gz43_215027	F	M00055862D:D06	UC2-NormColon
740_	648819	1695.K11.gz43_215033	F	M00055871B:B03	UC2-NormColon
741	644755	1695.L11.gz43_215034	F	M00055872A:C08	UC2-NormColon
742	516512	1695.P11.gz43_215038	F	M00055875C;C07	UC2-NormColon
743	380550	1695.L12.gz43_215050	F	M00055872A;D08	UC2-NormColon
744	645538	1695.D13.gz43_215058	F	M00055861C:G03	UC2-NormColon
745	644047	1695.F13.gz43_215060	F	M00055864A;C09	UC2-NormColon
746	650773	1695.F14.gz43_215076	F	M00055864A:E11	UC2-NormColon
747	449936	1695.J14.gz43_215080	F	M00055869C:G06	UC2-NormColon
748	554646	1695.O14.gz43_215085	F	M00055875A:G05	UC2-NormColon
749	640534	1695.L15.gz43_215098	F	M00055872A:E11	UC2-NormColon
750	642411	1695.J16.gz43_215112 1695.M16.gz43_215115	F	M00055869D:A07	UC2-NormColon
751 752	643451 644435	1695.H17.gz43_215126	F	M00055873A:E03 M00055866C:G09	UC2-NormColon UC2-NormColon
753	561877	1695.F18.gz43_215126	F	M00055864B;C09	UC2-NormColon
754	645497	1695.N18.gz43_215148	F	M00055874A:F06	UC2-NormColon
755	644030	1695.E20.gz43_215171	F	M00055863B:C07	UC2-NormColon
756	140224	1695.D21.gz43 215186	F	M00055862A:C01	UC2-NormColon
130	170224	1075.DZ1.gZ45_Z15160		11100033802A;C01	CC2-140IIICOIOII

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1	able 2				
SEQ			ODEN		
ID NO	CLUSTER	SEO NAME	ORIEN T	CLONE ID	TIDDADA
	CAUGITAN			CLONE ID	LIBRARY
757	643099	1695.K21.gz43_215193	F	M00055871C:C07	UC2-NormColon
758	643333	1695.P22.gz43_215214	F	M00055875D:D01	UC2-NormColon
759	649660	1695.G23.gz43_215221	F	M00055865C:G11	UC2-NormColon
760	645288	1695.H23.gz43_215222	F	M00055867A:B02	UC2-NormColon
761	650517	1695.G24.gz43_215237	F	M00055865C:H06	UC2-NormColon
762	552201	1696.J01.gz43_215256	F	M00055885C:B07	UC2-NormColon
763	643513	1696.P02.gz43_215278	F	M00055891B:F09	UC2-NormColon
764	447807	1696.D04.gz43_215298	F	M00055880B:A06	UC2-NormColon
765	639256	1696.M04.gz43_215307	F	M00055888C:F07	UC2-NormColon
766	179760	1696.G05.gz43_215317	F	M00055883A:C02	UC2-NormColon
767	459274	1696.G06.gz43_215333	F	M00055883A:C10	UC2-NormColon
768	345761	1696.I06.gz43_215335	F	M00055884D:F07	UC2-NormColon
769	640356	1696.M06.gz43_215339	F	M00055888C:G09	UC2-NormColon
770	645638	1696.A07.gz43_215343	F	M00055877A:H04	UC2-NormColon
771	464171	1696.N11.gz43_215420	F	M00055890A:A06	UC2-NormColon
772	446225	1696.E13.gz43_215443	F	M00055881A:G07	UC2-NormColon
773	450559	1696.F14.gz43_215460	F	M00055882C:A06	UC2-NormColon
774	643248	1696.L14.gz43 215466	F	M00055887D:C11	UC2-NormColon
775	549114	1696.N14.gz43 215468	F	M00055890A:D01	UC2-NormColon
776	645215	1696.D15.gz43 215474	F	M00055880C:F07	UC2-NormColon
777	555172	1696.I15.gz43 215479	F	M00055885B:A11	UC2-NormColon
778	593715	1696.B16.gz43 215488	F	M00055878C:C02	UC2-NormColon
779	586992	1696.I16.gz43 215495	F	M00055885B:B04	UC2-NormColon
780	451670	1696.H17.gz43 215510	F	M00055884B:F10	UC2-NormColon
781	552628	1696.H18.gz43 215526	F	M00055884B:H07	UC2-NormColon
782	643233	1696.A21.gz43_215567	F	M00055877D:C05	UC2-NormColon
783	640400	1696.F21.gz43 215572	F	M00055882D:B02	UC2-NormColon
784	644468	1696.G21.gz43 215573	F	M00055883D:B06	UC2-NormColon
785	643046	1696.H21.gz43 215574	F	M00055884C:B07	UC2-NormColon
786	649068	1696.N21.gz43_215580	F	M00055890A:G12	UC2-NormColon
787	648063	1696.E22.gz43 215587	F	M00055881D:A02	UC2-NormColon
788	570248	1696,M23,gz43 215611	F	M00055889B:E12	UC2-NormColon
789	643724	1696.N23.gz43 215612	F	M00055890A:H11	UC2-NormColon
790	649027	1696.P23.gz43_215614	F	M00055892A:F04	UC2-NormColon
791	643366	1696.P24.gz43 215630	F	M00055892B:D02	UC2-NormColon
792	526575	1705.E01.gz43_215635	F	M00055900D:D03	UC2-NormColon
793	449737	1705.Q02.gz43_215661	F	M00055914A:A03	UC2-NormColon
794	394373	1705.B03.gz43_215664	F	M00055894A:A03	UC2-NormColon
795	557361	1705.C03.gz43_215665	F	M00055894B:E09	UC2-NormColon
796		1705.K04.gz43_215689	F		UC2-NormColon
797	644364 644242		F	M00055908D:F09	UC2-NormColon
		1705.M05.gz43_215707		M00055911B:E06	
798	644407	1705.N05.gz43 215708	F	M00055912D:F04	UC2-NormColon

Table 2

	able 2		_		
SEO			ORIEN		
	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
799	643513	1705.F06.gz43 215716	F	M00055902A:H11	UC2-NormColon
800	639194	1705.H06.gz43 215718	F	M00055905B:H01	UC2-NormColon
801	454720	1705.I06.gz43 215719	F	M00055906D:G03	UC2-NormColon
802	643300	1705.009.gz43 215773	F	M00055914C:D12	UC2-NormColon
803	647522	1705.E10.gz43_215779	F	M00055901B;A02	UC2-NormColon
804	648576	1705.B11.gz43_215792	F	M00055896A:G01	UC2-NormColon
805	485237	1705.C11.gz43 215793	F	M00055898B:E07	UC2-NormColon
806	644108	1705.C11.gz43_215793	F	M00055914C:G01	UC2-NormColon
807	643914	1705.G12.gz43_215813	F	M00055904B:B01	UC2-NormColon
808	641645	1705.Q12.gz43_215821	F	M00055914C:G11	UC2-NormColon
809	605596	1705.H13.gz43 215830	F	M00055905D:B06	UC2-NormColon
810	564134	1705.O13.gz43_215837	F	M00055914D:A08	UC2-NormColon
811	643948	1705.B14.gz43_215840	F	M00055896B:C06	UC2-NormColon
812	557947	1705.F14.gz43_215844	F	M00055902D:A07	UC2-NormColon
813	15001	1705.G15.gz43_215861	F	M00055904C:A07	UC2-NormColon
814	649170	1705.P15.gz43_215870	F	M00055916B:C02	
815	644390	1705.D16.gz43_215874	F		UC2-NormColon
816	643089	1705.K16.gz43_215881	F	M00055900A:F08	UC2-NormColon
817	553587	1705.O16.gz43_215885	F	M00055909B:G10	UC2-NormColon
818	550242	1705.H17.gz43 215894	F	M00055914D:G06 M00055905D:H02	UC2-NormColon
819	639413	1705.O17.gz43_215894	F		UC2-NormColon
820	645844	1705.C18.gz43_215905	F	M00055915A:A06 M00055899A:B03	UC2-NormColon
821			F		UC2-NormColon
822	641087	1705.E18.gz43_215907 1705.F18.gz43_215908		M00055901C:C10	UC2-NormColon
823	482163 643808	1705.G18.gz43_215908	F	M00055902D:H10	UC2-NormColon
824		1705.L19.gz43_215930	F	M00055904D:A06	UC2-NormColon
825	468147 446728	1705.D20.gz43_215938	F	M00055910C:G01	UC2-NormColon
				M00055900B:E12	UC2-NormColon
826	447326	1705.K20.gz43_215945	F	M00055909C:E08	UC2-NormColon
827	685001	1705.L20.gz43_215946	F	M00055910C:G04	UC2-NormColon
828	645781	1705.K21.gz43_215961	F	M00055909C:G09	UC2-NormColon
829	643825	1705.L21.gz43_215962	F	M00055910D:A03	UC2-NormColon
830	374340	1705.A22.gz43_215967	F	M00055894A:H08	UC2-NormColon
831	640756	1705.M22.gz43_215979	F	M00055912C:C08	UC2-NormColon
832	400741	1705.B23.gz43_215984	F	M00055896C:H10	UC2-NormColon
833	644063	1705.F24.gz43_216004	F	M00055903B:C10	UC2-NormColon
834	644354	1705.G24.gz43_216005	F	M00055905A:F11	UC2-NormColon
835	89082	1705.K24.gz43_216009	F	M00055909D:E01	UC2-NormColon
836	643991	1705.024.gz43_216013	F	M00055915C:B09	UC2-NormColon
837	459881	1706.B01.gz43_216028	F	M00055918B:B10	UC2-NormColon
838	644554	1706.C01.gz43_216029	F	M00055919D:H07	UC2-NormColon
839	557175	1706.B03.gz43_216060	F	M00055918B:F10	UC2-NormColon
840	641484	1706.F03.gz43_216064	F	M00055924C:A09	UC2-NormColon

Table 2

844 446795 1706.M06.gza43.216119 F M00055934C:E06 UC2-NormColon 845 63808 1706.L07.gzd3.216134 F M00055933A:E05 UC2-NormColon 846 646596 1706.N07.gzd3.216136 F M000559398:A05 UC2-NormColon 847 639901 1706.N08.gzd3.216138 F M000559398:A01 UC2-NormColon 848 648379 1706.N08.gzd3.216164 F M000559300:D03 UC2-NormColon 849 650938 1706.109.gzd3.216164 F M00055923B:C07 UC2-NormColon 850 643170 1706.H10.gzd3.216178 F M00055927B:C01 UC2-NormColon 851 643239 1706.H10.gzd3.216180 F M00055930B:E00 UC2-NormColon 852 639596 1706.H10.gzd3.216180 F M00055993B:D30 UC2-NormColon 854 45590 1706.M11.gzd3.216200 F M00055991A:D60 UC2-NormColon 855 647952 1706.B11.gzd3.216200 F M00055991A:D60 UC2-NormColon		able 2				
Section	SEQ			ORIEN		
842 644629 1706.103 gz43 216100 F M00055930A:G04 UC2-NormColon 843 419009 1706.E06 gz43 216111 F M0005593CB:G06 UC2-NormColon 844 446795 1706.M06 gz43 216119 F M0005593CB:G06 UC2-NormColon 845 638808 1706.L07 gz43 216134 F M0005593A:E05 UC2-NormColon 846 646596 1706.N07 gz43 216138 F M0005593BA:10 UC2-NormColon 847 639901 1706.P07 gz43 216152 F M0005593BA:11 UC2-NormColon 848 648379 1706.N08 gz43 216164 F M0005593BA:11 UC2-NormColon 850 643170 1706.B10 gz43 216175 F M0005593BC:O103 UC2-NormColon 851 643239 1706.H10 gz43 216178 F M0005593BC:O100 UC2-NormColon 851 643239 1706.H10 gz43 216180 F M0005593BC:O100 UC2-NormColon 853 66114 1706.L10 gz43 21620 F M0005593BB:10 UC2-NormColon 854	ID NO	CLUSTER	SEQ NAME	Т	CLONE ID	LIBRARY
842 644629 1706,105,gz43, 216100 F M00055930A.G04 UC2-NormColon 843 41909 1706,E06,gz43, 216111 F M0005592D.G10 UC2-NormColon 844 446795 1706,M06,gz43, 216119 F M00055934C:E06 UC2-NormColon 845 63808 1706,L07,gz43, 216134 F M00055933A:E05 UC2-NormColon 846 646596 1706,N07,gz43, 216138 F M000559393C:B03 UC2-NormColon 847 639901 1706,N08,gz43, 216132 F M000559390C:D03 UC2-NormColon 848 648379 1706,N08,gz43, 216152 F M00055930C:D03 UC2-NormColon 850 643170 1706,E10,gz43, 216178 F M00055930C:D03 UC2-NormColon 851 643239 1706,H10,gz43, 216180 F M00055933B:C07 UC2-NormColon 852 639596 1706,H10,gz43, 216180 F M00055930C:P03 UC2-NormColon 853 605114 1706,L10,gz43, 216200 F M00055930C:P03 UC2-NormColon	841	651131	1706.B05.gz43 216092	F	M00055918C:C04	UC2-NormColon
843 419009 1706.B06.gz43 216111 F M00055922D.G10 UC2-NormColon 844 446795 1706.M06.gz43 216134 F M00055934C.E06 UC2-NormColon 845 638808 1706.L07.gz43 216134 F M00055933A.E05 UC2-NormColon 846 646596 1706.N07.gz43 216136 F M00055939B.A11 UC2-NormColon 847 639901 1706.N08.gz43 216152 F M00055939B.A11 UC2-NormColon 848 648379 1706.N08.gz43 216152 F M00055930C.D03 UC2-NormColon 850 643170 1706.E10.gz43 216178 F M00055930C.D03 UC2-NormColon 851 643239 1706.H10.gz43 216178 F M0005593TD.E11 UC2-NormColon 852 639596 1706.H10.gz43 216180 F M0005593B.B11 UC2-NormColon 853 605114 1706.110.gz43 216200 F M0005593B.B11 UC2-NormColon 854 456920 1706.N11.gz43 216200 F M0005593B.D103 UC2-NormColon 8	842	644629	1706.J05.gz43_216100	F	M00055930A:G04	
845 638808 1706 L07 gz43 216134 F M00055933A:E05 UC2-NormColon 846 646596 1706 NO7 gz43 216136 F M00055936:B05 UC2-NormColon 847 639901 1706 P07 gz43 216138 F M00055938A:11 UC2-NormColon 848 648379 1706.N08 gz43 216152 F M00055938A:11 UC2-NormColon 849 650938 1706.N08 gz43 216164 F M00055938L07 UC2-NormColon 850 643170 1706.B10 gz43 216175 F M00055938L07 UC2-NormColon 851 643239 1706.H10 gz43 216178 F M00055938L07 UC2-NormColon 852 639596 1706.H10 gz43 216180 F M00055938L07 UC2-NormColon 852 639596 1706.H10 gz43 21620 F M00055938B11 UC2-NormColon 853 605114 1706.L10 gz43 21620 F M0005593B110 UC2-NormColon 854 456920 1706.N11 gz43 21620 F M0005593B1400 UC2-NormColon 855 <	843	419009	1706.E06.gz43 216111	F	M00055922D:G10	UC2-NormColon
846 646596 1706.N07.gz43 216136 F M00055936C:B05 UC2-NormColon 847 639901 1706.P07.gz43 216138 F M00055938:A11 UC2-NormColon 848 648379 1706.N08.gz43 216152 F M00055930:D03 UC2-NormColon 849 650938 1706.09.gz43 216164 F M00055930:D03 UC2-NormColon 850 643170 1706.E10.gz43 216178 F M00055923B:C07 UC2-NormColon 851 643239 1706.H10.gz43 216178 F M00055927B:C07 UC2-NormColon 852 639596 1706.H10.gz43 216180 F M00055930C:P06 UC2-NormColon 853 605114 1706.L10.gz43 216200 F M00055933B:B10 UC2-NormColon 854 456920 1706.B11.gz43 216200 F M00055934D:H09 UC2-NormColon 855 647952 1706.B12.gz43 216224 F M00055934D:H09 UC2-NormColon 857 465576 1706.D13.gz43 216225 F M00055934D:H09 UC2-NormColon 859	844	446795	1706.M06.gz43 216119	F	M00055934C:E06	UC2-NormColon
847 639901 1706.P07_gz43_216138 F M00055939E:A11 UC2-NormColon 848 648379 1706.N08_gz43_216152 F M00055936:CD03 UC2-NormColon 849 650938 1706.008_gz43_216164 F M00055930:CD03 UC2-NormColon 850 643170 1706.E10_gz43_216175 F M00055902D:C07 UC2-NormColon 851 643239 1706.H10_gz43_216180 F M00055902TD:E11 UC2-NormColon 852 6399596 1706.H10_gz43_216180 F M00055903D:F06 UC2-NormColon 853 605114 1706.L10_gz43_216180 F M0005593B:D13 UC2-NormColon 854 456920 1706.N11_gz43_216204 F M0005593AD:F03 UC2-NormColon 855 647952 1706.B11_gz43_216204 F M00055993D:H03 UC2-NormColon 857 465576 1706.M12_gz43_216225 F M0005593D:D07 UC2-NormColon 857 465576 1706.B13_gz43_216224 F M0005593D:D07 UC2-NormColon	845	638808	1706.L07.gz43_216134	F	M00055933A:E05	UC2-NormColon
848 648379 1706.N08.gx43_216162 F M00055930C:D03 UC2-NormColon 849 650938 1706.10g.gx43_216164 F M00055930C:D03 UC2-NormColon 850 643170 1706.E10.gx43_216175 F M0005593B.C7 UC2-NormColon 851 643239 1706.H10.gx43_21618 F M0005593B.C7 UC2-NormColon 852 639596 1706.H10.gx43_216182 F M0005593B.B11 UC2-NormColon 853 605114 1706.H10.gx43_216182 F M0005593B.B11 UC2-NormColon 854 456920 1706.N11.gx43_216200 F M0005593B.B11 UC2-NormColon 855 647952 1706.B11.gx43_216215 F M0005593H.D90 UC2-NormColon 856 650470 1706.M10.gx43_216215 F M0005593H.D90 UC2-NormColon 857 465576 1706.D13.gx43_216222 F M0005593B.A109 UC2-NormColon 858 455413 1706.D13.gx43_216255 F M0005593B.A26 UC2-NormColon 859	846	646596	1706.N07.gz43_216136	F	M00055936C:B05	UC2-NormColon
849 650938 1706.09 gz43 216164 F M00055930C:D03 UC2-NormColon 850 643170 1706.E10 gz43 216175 F M00055923B:C07 UC2-NormColon 851 643239 1706.B10 gz43 216178 F M00055927B:C07 UC2-NormColon 852 639596 1706.D10 gz43 216180 F M00055930C:P06 UC2-NormColon 853 605114 1706.L10 gz43 216180 F M00055933B:B11 UC2-NormColon 854 456920 1706.N11 gz43 216200 F M00055939B:D673 UC2-NormColon 855 647952 1706.B12 gz43 216204 F M00055934D:H09 UC2-NormColon 856 650470 1706.B12 gz43 216225 F M00055934D:H09 UC2-NormColon 857 465576 1706.D13 gz43 216224 F M00055934D:H09 UC2-NormColon 859 639038 1706.H15 gz43 216256 F M00055930D:D07 UC2-NormColon 860 556925 1706.H15 gz43 216260 F M00055930D:B05 UC2-NormColon <td< td=""><td>847</td><td>639901</td><td>1706.P07.gz43_216138</td><td>F</td><td>M00055939B:A11</td><td>UC2-NormColon</td></td<>	847	639901	1706.P07.gz43_216138	F	M00055939B:A11	UC2-NormColon
850 643170 1706.E10.gz43_216175 F M00055923B:C07 UC2-NormColon 851 643239 1706.H10.gz43_216180 F M00055992D:E11 UC2-NormColon 852 639596 1706.H10.gz43_216180 F M00055993C:P66 UC2-NormColon 853 605114 1706.L10.gz43_216182 F M0005593B:B11 UC2-NormColon 854 4356920 1706.N11.gz43_216200 F M00055993D:P03 UC2-NormColon 855 647952 1706.B11.gz43_216204 F M00055994D:H09 UC2-NormColon 856 650470 1706.M12.gz43_216215 F M00055994D:H09 UC2-NormColon 857 465576 1706.D13.gz43_216222 F M00055993D:D0 UC2-NormColon 858 45511_3 1706.113.gz43_216224 F M00055993D:D0 UC2-NormColon 859 639038 1706.H13.gz43_216260 F M00055993D:D0 UC2-NormColon 860 556925 1706.H15.gz43_216267 F M00055993D:D0 UC2-NormColon 86	848	648379	1706.N08.gz43_216152	F	M00055936C:D03	UC2-NormColon
851 643239 1706.H10.gz43_216178 F M00055927D:E11 UC2-NormColon 852 639596 1706.H10.gz43_216180 F M00055930C:P06 UC2-NormColon 833 605114 1706.110.gz43_216182 F M0005593BB11 UC2-NormColon 854 456920 1706.N11.gz43_216200 F M0005593BB11 UC2-NormColon 855 647952 1706.B12.gz43_216204 F M00055919A:A06 UC2-NormColon 856 650470 1706.M12.gz43_216215 F M0005594DH09 UC2-NormColon 857 465576 1706.D13.gz43_216222 F M0005593DDD07 UC2-NormColon 858 455413 1706.P13.gz43_216258 F M0005593AC06 UC2-NormColon 859 639038 1706.H15.gz43_216260 F M0005593AC06 UC2-NormColon 860 556925 1706.J15.gz43_216260 F M0005593DDD07 UC2-NormColon 861 644301 1706.A16.gz43_216267 F M0005593DDD05 UC2-NormColon 862	849	650938	1706.J09.gz43_216164	F	M00055930C:D03	UC2-NormColon
852 639596 1706,110,gz43_216180 F M00055930C:P06 UC2-NormColon 853 405114 1706,110,gz43_216182 F M00055933B:B11 UC2-NormColon 854 456920 1706,N11,gz43_216209 F M00055930B:P03 UC2-NormColon 855 647952 1706,B12,gz43_216210 F M0005593D:A:A06 UC2-NormColon 856 650470 1706,M12,gz43_216215 F M0005593D:A:A06 UC2-NormColon 857 465576 1706,D13,gz43_216222 F M0005593D:D:D07 UC2-NormColon 858 455413 1706,P13,gz43_216234 F M0005593D:D:D07 UC2-NormColon 859 639038 1706,H15,gz43_216260 F M0005593D:D:D07 UC2-NormColon 860 556925 1706,115,gz43_216260 F M0005593D:D:D07 UC2-NormColon 861 644301 1706,116,gz43_216267 F M0005593D:D:D07 UC2-NormColon 862 4477520 1706,117,gz43_216286 F M0005593D:D:D07 UC2-NormColon	850	643170	1706.E10.gz43_216175	F	M00055923B:C07	UC2-NormColon
853 605114 1706.1.10 gz43 216182 F M00055933B:B11 UC2-NormColon 854 456920 1706.N11 gz43 216200 F M00055936D:F03 UC2-NormColon 855 649521 1706.B11 gz43 216204 F M00055993A:A06 UC2-NormColon 856 650470 1706.M12 gz43 216215 F M00055993A:DH09 UC2-NormColon 857 465576 1706.D13 gz43 216222 F M00055993D:D07 UC2-NormColon 858 455413 1706.P13 gz43 216234 F M0005599D:D07 UC2-NormColon 859 639038 1706.H15 gz43 216260 F M0005599D:D07 UC2-NormColon 860 556925 1706.J15 gz43 216260 F M00055990:D05 UC2-NormColon 861 644301 1706.A16 gz43 216267 F M0005591D:E07 UC2-NormColon 862 447520 1706.J16 gz43 216266 F M0005591D:E07 UC2-NormColon 864 610893 1706.H17 gz43 216290 F M0005593A:E11 UC2-NormColon 865	851	643239	1706.H10.gz43_216178	F	M00055927D:E11	UC2-NormColon
854 456920 1706.N11.gz43_216200 F M00055936D:P03 UC2-NormColon 855 647952 1706.B12.gz43_216204 F M00055939A:D00 UC2-NormColon 856 650470 1706.M12.gz43_216215 F M0005593DA:D00 UC2-NormColon 857 465576 1706.D13.gz43_216222 F M0005593D:D07 UC2-NormColon 858 455413 1706.P13.gz43_216234 F M0005593A:C06 UC2-NormColon 859 639038 1706.B115.gz43_216260 F M0005593A:C06 UC2-NormColon 860 556925 1706.J15.gz43_216260 F M0005593DC:H05 UC2-NormColon 861 644301 1706.A16.gz43_216267 F M0005593DB:D0 UC2-NormColon 862 447520 1706.B11.gz43_216266 F M0005593DB:D0 UC2-NormColon 864 467520 1706.B11.gz43_216266 F M0005593DB:D0 UC2-NormColon 864 467520 1706.B11.gz43_216296 F M0005593DB:D0 UC2-NormColon 865 </td <td>852</td> <td>639596</td> <td>1706.J10.gz43_216180</td> <td>F</td> <td>M00055930C:F06</td> <td>UC2-NormColon</td>	852	639596	1706.J10.gz43_216180	F	M00055930C:F06	UC2-NormColon
855 647952 1706.B12.gz43_216204 F M00055919A:A06 UC2-NormColon 856 650470 1706.M12.gz43_216215 F M00055934D:H09 UC2-NormColon 857 465576 1706.D13.gz43_216222 F M00055924A:P05 UC2-NormColon 858 455413 1706.P13.gz43_216234 F M00055928A:C06 UC2-NormColon 859 639038 1706.P13.gz43_216260 F M00055928A:C06 UC2-NormColon 860 556925 1706.J15.gz43_216260 F M0005592A:D15 UC2-NormColon 861 644301 1706.J16.gz43_216267 F M00055930D:B05 UC2-NormColon 862 447520 1706.J16.gz43_216286 F M00055930D:B05 UC2-NormColon 864 610893 1706.B17.gz43_216296 F M0005592B:A:E11 UC2-NormColon 865 613626 1705.N17.gz43_216296 F M00055937A:F07 UC2-NormColon 866 645092 1706.N17.gz43_216236 F M00055937A:F07 UC2-NormColon <th< td=""><td>853</td><td>605114</td><td>1706.L10.gz43_216182</td><td>F</td><td>M00055933B:B11</td><td>UC2-NormColon</td></th<>	853	605114	1706.L10.gz43_216182	F	M00055933B:B11	UC2-NormColon
856 650470 1706.M12_gz43_216215 F M00055934D:H09 UC2-NormColon 857 465576 1706.D13_gz43_216222 F M00055993D:D07 UC2-NormColon 858 455413 1706.P13_gz43_216234 F M0005599D:D07 UC2-NormColon 859 639038 1706.H15_gz43_216268 F M0005599D:D07 UC2-NormColon 860 556925 1706.J15_gz43_216260 F M00055990C:H05 UC2-NormColon 861 644301 1706.A16_gz43_216267 F M0005591D:E07 UC2-NormColon 862 447520 1706.J16_gz43_216266 F M0005592B:A06 UC2-NormColon 864 61093 1706.H17_gz43_216290 F M0005592B:A06 UC2-NormColon 865 613626 1706.J17_gz43_216290 F M0005593AB:E11 UC2-NormColon 866 645092 1706.N17_gz43_216290 F M0005593AB:E11 UC2-NormColon 867 649873 1706.P19_gz43_216330 F M0005593AB:A81 UC2-NormColon 868 </td <td>854</td> <td>456920</td> <td>1706.N11.gz43_216200</td> <td>F</td> <td>M00055936D:F03</td> <td>UC2-NormColon</td>	854	456920	1706.N11.gz43_216200	F	M00055936D:F03	UC2-NormColon
857 465576 1706.D13 gz43 216222 F M00055922A:F05 UC2-NormColon 858 455413 1706.P13 gz43 216234 F M0005593A:C0 UC2-NormColon 859 639038 1706.P13 gz43 216238 F M0005593A:C0 UC2-NormColon 860 556925 1706.J15 gz43 216260 F M0005593C:H05 UC2-NormColon 861 644301 1706.A16 gz43 216267 F M0005593D:D10-D7 UC2-NormColon 862 447520 1706.D16 gz43 216276 F M0005593D:D05 UC2-NormColon 863 447426 1706.D17 gz43 216296 F M0005593D:B05 UC2-NormColon 864 610893 1706.H17 gz43 216290 F M0005593D:B05 UC2-NormColon 865 613626 1706.D17 gz43 216290 F M0005593D:B09 UC2-NormColon 866 645092 1706.N17 gz43 216390 F M0005594D:A08 UC2-NormColon 867 649871 1706.P21 gz43 216331 F M0005594D:A08 UC2-NormColon 870	855	647952	1706.B12.gz43_216204	F	M00055919A:A06	UC2-NormColon
858 455413 1706.P13.gz43_216234 F M00055939D:D07 UC2-NormColon 859 639038 1706.H15.gz43_216238 F M0005593CH.00 UC2-NormColon 860 556952 1706.H15.gz43_216260 F M0005593CH.05 UC2-NormColon 861 644301 1706.A16.gz43_216267 F M0005593D:DE07 UC2-NormColon 862 447520 1706.H16.gz43_216236 F M0005593D:DB05 UC2-NormColon 863 447426 1706.D17.gz43_216286 F M0005593AB:A66 UC2-NormColon 864 610893 1706.H17.gz43_216290 F M0005593AB:A61 UC2-NormColon 865 613626 1706.H17.gz43_216296 F M00055937A:F07 UC2-NormColon 866 645092 1706.N17.gz43_216296 F M00055937A:F07 UC2-NormColon 867 649873 1706.N17.gz43_216330 F M00055937A:F07 UC2-NormColon 867 649803 1706.D21.gz43_216330 F M0005592B:D09 UC2-NormColon 87	856	650470	1706.M12.gz43_216215	F	M00055934D:H09	UC2-NormColon
859 639038 1706.H15.gz43_216258 F M00055928A:C06 UC2-NormColon 860 556925 1706.J15.gz43_216260 F M00055930C:H05 UC2-NormColon 861 644301 1706.A16.gz43_216267 F M0005591D:LDG UC2-NormColon 862 447520 1706.J16.gz43_216266 F M00055930D:B05 UC2-NormColon 863 447426 1706.D17.gz43_216286 F M0005592B:A06 UC2-NormColon 864 610893 1706.H17.gz43_216290 F M0005593AE.II UC2-NormColon 865 613626 1706.J17.gz43_216290 F M0005593D:B09 UC2-NormColon 866 645092 1706.N17.gz43_216296 F M0005593TA:F07 UC2-NormColon 867 649873 1706.P19.gz43_216330 F M0005594BA:08 UC2-NormColon 868 645028 1706.A20.gz43_216331 F M0005594BA:08 UC2-NormColon 870 649085 1706.D21.gz43_216350 F M00055928B:D10 UC2-NormColon 871 </td <td>857</td> <td>465576</td> <td>1706.D13.gz43_216222</td> <td>F</td> <td>M00055922A:F05</td> <td>UC2-NormColon</td>	857	465576	1706.D13.gz43_216222	F	M00055922A:F05	UC2-NormColon
860 556925 1706,115,gz43, 216260 F M00055930C:H05 UC2-NormColon 861 644301 1706,A16,gz43, 216267 F M00055930D:B05 UC2-NormColon 862 447520 1706,B16,gz43, 216276 F M00055930D:B05 UC2-NormColon 863 447426 1706,D17,gz43, 216286 F M0005592B:A06 UC2-NormColon 864 610893 1706,H17,gz43, 216290 F M0005592B:A01 UC2-NormColon 865 613626 1706,D17,gz43, 216292 F M00055937A:E11 UC2-NormColon 866 645092 1706,N17,gz43, 216396 F M00055940B:A08 UC2-NormColon 867 649871 1706,P19,gz43, 216330 F M00055940B:A08 UC2-NormColon 869 639154 1706,P21,gz43, 216330 F M0005592B:D09 UC2-NormColon 870 649085 1706,D21,gz43, 216352 F M0005592B:D09 UC2-NormColon 871 643332 1706,F21,gz43, 216352 F M0005592B:D10 UC2-NormColon	858	455413	1706.P13.gz43_216234	F	M00055939D:D07	UC2-NormColon
861 644301 1706.A16 gz43 216267 F M00055917D.E07 UC2-NormColon 862 447520 1706.J16 gz43 216276 F M00055930D.B05 UC2-NormColon 863 447426 1706.D17 gz43 216286 F M00055923B.A06 UC2-NormColon 864 610893 1706.H17 gz43 216290 F M00055923A.E11 UC2-NormColon 865 613626 1706.H17 gz43 216296 F M00055937A.F07 UC2-NormColon 866 645092 1706.N17 gz43 216296 F M00055937A.F07 UC2-NormColon 867 649873 1706.P19 gz43 216330 F M00055940B.A08 UC2-NormColon 868 645028 1706.D21 gz43 216330 F M0005592B.D09 UC2-NormColon 870 649085 1706.D21 gz43 216356 F M0005592B.D09 UC2-NormColon 871 643332 1706.P12 gz43 216355 F M0005592B.D09 UC2-NormColon 872 643333 1706.H21 gz43 216354 F M0005592B.D10 UC2-NormColon 873	859	639038	1706.H15.gz43_216258	F	M00055928A:C06	UC2-NormColon
862 447520 1706.116.gz43_216276 F M00055930D:B05 UC2-NormColon 863 447426 1706.D17.gz43_216286 F M00055932B:A06 UC2-NormColon 864 61093 1706.B17.gz43_216290 F M00055932B:A06 UC2-NormColon 865 613626 1706.J17.gz43_216292 F M00055930D:B09 UC2-NormColon 866 645092 1706.N17.gz43_216292 F M00055937A:F07 UC2-NormColon 867 649873 1706.P19.gz43_216330 F M00055994B:A08 UC2-NormColon 869 639154 1706.P20.gz43_216330 F M0005592B:D09 UC2-NormColon 870 649085 1706.D21.gz43_216350 F M0005592B:D09 UC2-NormColon 871 643332 1706.P21.gz43_216350 F M0005592B:D10 UC2-NormColon 872 649085 1706.D21.gz43_216350 F M0005592B:B1D UC2-NormColon 873 204 1706.J21.gz43_216350 F M0005592B:B1D UC2-NormColon 874	860	556925	1706.J15.gz43_216260	F	M00055930C:H05	UC2-NormColon
863 447426 1706.D17_gz43_216286 F M00055922B:A06 UC2-NormColon 864 610893 1706.H17_gz43_216290 F M00055992A:E11 UC2-NormColon 865 613626 1706.H17_gz43_216292 F M000559930.B09 UC2-NormColon 866 645092 1706.N17_gz43_216230 F M000559910.B09 UC2-NormColon 867 649873 1706.P19_gz43_216330 F M00055918.H0 UC2-NormColon 868 645028 1706.D20_gz43_216331 F M00055918.H0 UC2-NormColon 870 649085 1706.D21_gz43_216330 F M0005592B:D09 UC2-NormColon 871 643332 1706.P21_gz43_216330 F M0005592B:D09 UC2-NormColon 872 643333 1706.P21_gz43_216350 F M0005592B:D10 UC2-NormColon 873 204 1706.P21_gz43_216354 F M0005593E:D10 UC2-NormColon 874 641716 1706.N21_gz43_216360 F M000559924B:D10 UC2-NormColon 875	861	644301	1706.A16.gz43_216267	F	M00055917D:E07	UC2-NormColon
864 610893 1706.H17_gz43_216290 F M00055928A:E11 UC2-NormColon 865 613626 1706.H17_gz43_216292 F M00055930D:B09 UC2-NormColon 866 645092 1706.N17_gz43_216295 F M0005594D:BA08 UC2-NormColon 867 649873 1706.P19_gz43_216330 F M0005594B:A08 UC2-NormColon 868 645028 1706.A20_gz43_216331 F M0005592B:D09 UC2-NormColon 869 639154 1706.P20_gz43_216350 F M0005592B:D09 UC2-NormColon 870 649085 1706.P21_gz43_216350 F M0005592B:D09 UC2-NormColon 871 643332 1706.P21_gz43_216352 F M0005592B:D10 UC2-NormColon 872 643383 1706.H21_gz43_216355 F M0005592B:D10 UC2-NormColon 873 204 1706.E12_gz43_216356 F M0005592B:D05 UC2-NormColon 874 641716 1706.N21_gz43_216360 F M00055992B:D02 UC2-NormColon 875	862	447520	1706.J16.gz43_216276	F	M00055930D:B05	UC2-NormColon
865 613626 1706,117,gx43_216292 F M00055930D:B09 UC2-NormColon 866 64592 1706,N17,gx43_216296 F M00055991A:F07 UC2-NormColon 867 649873 1706,P19,gx43_216330 F M00055994B:A08 UC2-NormColon 868 645028 1706,A20,gx43_216331 F M00055918A:F10 UC2-NormColon 869 639154 1706,P20,gx43_216356 F M0005592B:D09 UC2-NormColon 870 649085 1706,D21,gx43_216350 F M0005592B:D09 UC2-NormColon 871 643332 1706,P21,gx43_216352 F M0005592B:D10 UC2-NormColon 872 6493332 1706,P21,gx43_216354 F M0005592B:B11 UC2-NormColon 873 204 1706,D21,gx43_216356 F M00055937C:C08 UC2-NormColon 874 641716 1706,N21,gx43_216360 F M00055937C:C08 UC2-NormColon 875 643627 1706,M23,gx43_216319 F M000559935C:E03 UC2-NormColon 877 </td <td>863</td> <td>447426</td> <td>1706.D17.gz43_216286</td> <td>F</td> <td>M00055922B:A06</td> <td>UC2-NormColon</td>	863	447426	1706.D17.gz43_216286	F	M00055922B:A06	UC2-NormColon
866 645092 1706.N17_gz43_216296 F M00055937A.F07 UC2-NormColon 867 649873 1706.P19_gz43_216330 F M00055940B:A08 UC2-NormColon 868 645028 1706.A20_gz43_216331 F M00055918A;F10 UC2-NormColon 869 639154 1706.P20_gz43_216336 F M0005592B;D09 UC2-NormColon 870 649085 1706.D21_gz43_216350 F M0005592B;D09 UC2-NormColon 871 643332 1706.F121_gz43_216352 F M0005592B;D10 UC2-NormColon 872 643383 1706.H21_gz43_216354 F M0005592B;D10 UC2-NormColon 873 204 1706.J21_gz43_216356 F M00055927C08 UC2-NormColon 874 641716 1706.M21_gz43_216360 F M00055927C08 UC2-NormColon 875 643627 1706.M23_gz43_2164367 F M00055993C08 UC2-NormColon 876 640672 1706.M23_gz43_216433 F M000559950CB0 UC2-NormColon 878	864	610893	1706.H17.gz43_216290	F	M00055928A:E11	UC2-NormColon
867 649873 1706.P19.gz43_216330 F M00055940B:A08 UC2-NormColon 868 645028 1706.A20.gz43_216331 F M00055918A:F10 UC2-NormColon 869 639154 1706.P20.gz43_216336 F M00055923B:D09 UC2-NormColon 870 649085 1706.D21.gz43_216350 F M00055922B:G09 UC2-NormColon 871 643332 1706.P21.gz43_216352 F M00055928B:D10 UC2-NormColon 873 204 1706.P21.gz43_216356 F M00055928B:D10 UC2-NormColon 874 641716 1706.N21.gz43_216356 F M00055937C:C08 UC2-NormColon 875 643627 1706.B22.gz43_216360 F M00055924B:D02 UC2-NormColon 876 640672 1706.W23_gz43_216319 F M00055932E30 UC2-NormColon 877 550237 1707.B04.gz43_216453 F M00055954B:B10 UC2-NormColon 878 646317 1707.B04.gz43_216502 F M000559594B:B10 UC2-NormColon 879	865	613626	1706.J17.gz43_216292	F	M00055930D:B09	UC2-NormColon
868 645028 1706.A20_gz43_216331 F M00055918A:F10 UC2-NormColon 869 639154 1706.P20_gz43_216336 F M00055928:D09 UC2-NormColon 870 649085 1706.D21_gz43_216350 F M00055928:D09 UC2-NormColon 871 643332 1706.P21_gz43_216352 F M00055928:D10 UC2-NormColon 872 643333 1706.P21_gz43_216354 F M00055928:B111 UC2-NormColon 873 204 1706.121_gz43_216356 F M00055990-P05 UC2-NormColon 874 641716 1706.N21_gz43_216360 F M000559937C:C08 UC2-NormColon 875 643627 1706.M23_gz43_216367 F M00055993C:E03 UC2-NormColon 876 640672 1706.M23_gz43_216433 F M000559950:B01 UC2-NormColon 877 550237 1707.902_gz43_216433 F M000559942B:H10 UC2-NormColon 878 646317 1707.B04_gz43_216502 F M000559958C:E03 UC2-NormColon 880	866	645092	1706.N17.gz43_216296	F	M00055937A:F07	UC2-NormColon
869 639154 1706 F20 gz43 216336 F M00055925B:D09 UC2-NormColon 870 649085 1706.D21 gz43 216350 F M0005592B:G09 UC2-NormColon 871 643332 1706 F21 gz43 216352 F M0005592B:D10 UC2-NormColon 872 643383 1706 H21 gz43 216354 F M00055928B:E11 UC2-NormColon 873 204 1706.J21 gz43 216356 F M00055937C:08 UC2-NormColon 874 641716 1706.N21 gz43 216360 F M00055997C:08 UC2-NormColon 875 643627 1706.N21 gz43 216367 F M00055995C:B0 UC2-NormColon 876 640672 1706.M23 gz43 216431 F M00055995C:B0 UC2-NormColon 877 550237 1707.102 gz43 216435 F M00055995D:B0 UC2-NormColon 878 646317 1707.B04 gz43 216502 F M00055995C:B0 UC2-NormColon 880 649549 1707.F05 gz43 216513 F M000559958C:E03 UC2-NormColon	867	649873	1706.P19.gz43_216330	F	M00055940B:A08	UC2-NormColon
870 649085 1706.D21_gz43_216350 F M00055922B:G09 UC2-NormColon 871 643332 1706.F21_gz43_216352 F M00055928B:D10 UC2-NormColon 872 643333 1706.F21_gz43_216354 F M00055928B:D10 UC2-NormColon 873 204 1706.D21_gz43_216356 F M00055930D:P05 UC2-NormColon 874 641716 1706.B21_gz43_216360 F M00055993D:C08 UC2-NormColon 875 643627 1706.B22_gz43_216391 F M00055993B:D0 UC2-NormColon 876 640672 1706.M23_gz43_216391 F M0005593C:E03 UC2-NormColon 877 550237 1707.B04_gz43_216453 F M00055942B:H10 UC2-NormColon 878 646317 1707.B04_gz43_216502 F M000559942B:H10 UC2-NormColon 879 643968 1707.R05_gz43_216513 F M00055998C:E03 UC2-NormColon 880 649549 1707.F06_gz43_216513 F M000559942B:H10 UC2-NormColon	868	645028	1706.A20.gz43_216331	F	M00055918A:F10	UC2-NormColon
871 643332 1706.F21.gz43 216352 F M00055925B:D10 UC2-NormColon 872 643383 1706.H21.gz43 216354 F M00055928B:B11 UC2-NormColon 873 204 1706.D21.gz43 216356 F M00055990.P05 UC2-NormColon 874 641716 1706.N21.gz43 216360 F M000559937C:C08 UC2-NormColon 875 643627 1706.B22.gz43 216367 F M00055993C:E03 UC2-NormColon 876 640672 1706.M23.gz43 2163191 F M00055995C:B03 UC2-NormColon 877 550237 1707.B04.gz43 216453 F M00055994B:D10 UC2-NormColon 878 646317 1707.B04.gz43 216512 F M000559958C:E03 UC2-NormColon 879 643968 1707.F05.gz43 216513 F M00055998C:E03 UC2-NormColon 880 649549 1707.F05.gz43 216513 F M00055994D:B07 UC2-NormColon	869	639154	1706.F20.gz43_216336	F	M00055925B:D09	UC2-NormColon
872 643383 1706.H21.gz43_216354 F M00055928B:E11 UC2-NormColon 873 204 1706.J21.gz43_216356 F M00055930D:P05 UC2-NormColon 874 641716 1706.N21.gz43_216369 F M000559937C08 UC2-NormColon 875 643627 1706.E22.gz43_216367 F M00055993CE03 UC2-NormColon 876 640672 1706.M23.gz43_21633 F M00055995B:D10 UC2-NormColon 877 550237 1707.02.gz43_216433 F M00055994B:H10 UC2-NormColon 878 646317 1707.B04.gz43_216477 F M00055994B:H10 UC2-NormColon 879 643968 1707.F05.gz43_216513 F M00055998C:E03 UC2-NormColon 880 649549 1707.F06.gz43_216513 F M00055994D:B07 UC2-NormColon	870	649085	1706.D21.gz43_216350	F	M00055922B:G09	UC2-NormColon
873 204 1706.121_gz43_216356 F M00055930D:P05 UC2-NormColon 874 641716 1706.N21_gz43_216360 F M00055937C:C08 UC2-NormColon 875 643627 1706.E22_gz43_216367 F M00055924B:D02 UC2-NormColon 876 640672 1706.M23_gz43_216391 F M00055933C:E03 UC2-NormColon 877 550237 1707.102_gz43_216453 F M00055942B:H10 UC2-NormColon 878 646317 1707.B04_gz43_216477 F M00055942B:H10 UC2-NormColon 879 64398 1707.K05_gz43_216502 F M0005598C:E03 UC2-NormColon 880 649549 1707.F06_gz43_216513 F M00055942B:H10 UC2-NormColon	871	643332	1706.F21.gz43_216352	F	M00055925B:D10	UC2-NormColon
874 641716 1706.N21_gz43_216360 F M00055937C:C08 UC2-NormColon 875 643627 1706.B22_gz43_216367 F M000559524B:D02 UC2-NormColon 876 640672 1706.M23_gz43_216391 F M00055935:E03 UC2-NormColon 877 550237 1707.J02_gz43_216453 F M00055995:E03 UC2-NormColon 878 646317 1707.B04_gz43_216477 F M00055994B:H10 UC2-NormColon 879 643968 1707.F05_gz43_216513 F M00055994D:B07 UC2-NormColon 880 649549 1707.F05_gz43_216513 F M00055994D:B07 UC2-NormColon	872	643383		F	M00055928B:E11	UC2-NormColon
875 643627 1706.B22.gz43_216367 F M00055924B:D02 UC2-NormColon 876 640672 1706.M23.gz43_216391 F M00055935C:E03 UC2-NormColon 877 550237 1707.J02.gz43_216453 F M00055956D:B01 UC2-NormColon 878 646317 1707.B04.gz43_216477 F M0005594B:B10 UC2-NormColon 879 643968 1707.K05.gz43_216502 F M00055983C:E03 UC2-NormColon 880 649549 1707.F06.gz43_216513 F M0005594D:B07 UC2-NormColon	873	204	1706.J21.gz43_216356	F	M00055930D:F05	UC2-NormColon
876 640672 1706.M23.gz43 216391 F M00055935C:E03 UC2-NormColon 877 550237 1707.D02.gz43 216453 F M00055956D:B01 UC2-NormColon 878 646317 1707.B04.gz43 216477 F M00055942B:H10 UC2-NormColon 879 643968 1707.K05.gz43 216502 F M00055949D:B07 UC2-NormColon 880 649549 1707.F06.gz43 216513 F M00055949D:B07 UC2-NormColon		641716	1706.N21.gz43_216360	F	M00055937C:C08	UC2-NormColon
877 550237 1707.J02.gz43_216453 F M00055956D:B01 UC2-NormColon 878 646317 1707.B04.gz43_216477 F M00055942B:H10 UC2-NormColon 879 643968 1707.K05.gz43_216502 F M00055958C:E03 UC2-NormColon 880 649549 1707.F06.gz43_216513 F M00055949D:B07 UC2-NormColon	875	643627	1706.E22.gz43_216367	F	M00055924B:D02	UC2-NormColon
878 646317 1707.B04gz43 216477 F M00055942B:H10 UC2-NormColon 879 643968 1707.K05gz43 216502 F M00055958C:E03 UC2-NormColon 880 649549 1707.F06gz43 216513 F M0005594D:B07 UC2-NormColon		640672	1706,M23.gz43_216391	F	M00055935C:E03	UC2-NormColon
879 643968 1707.K05.gz43 216502 F M00055958C:E03 UC2-NormColon 880 649549 1707.F06.gz43 216513 F M00055949D:B07 UC2-NormColon		550237		F		UC2-NormColon
880 649549 1707.F06.gz43_216513 F M00055949D:B07 UC2-NormColon		646317	1707.B04.gz43_216477	F	M00055942B:H10	UC2-NormColon
	879			F	M00055958C:E03	UC2-NormColon
881 513610 1707 007 gr/3 216538 F M00055067A-E11 UC2 NormColon	880	649549		F	M00055949D:B07	UC2-NormColon
	881	513619	1707.O07.gz43_216538	F	M00055967A:F11	UC2-NormColon
882 460245 1707.I08.gz43_216548 F M00055955B:C06 UC2-NormColon	882	460245	1707.I08.gz43_216548	F	M00055955B:C06	UC2-NormColon

Table 2

	able 2				
SEQ			ORIEN	,	
ID NO	CLUSTER	SEQ NAME	Т	CLONE ID	LIBRARY
883	646061	1707.008.gz43_216554	F	M00055967A:G01	UC2-NormColon
884	645273	1707.009.gz43 216570	F	M00055967B:B12	UC2-NormColon
885	524546	1707.A12.gz43 216604	F	M00055941B:A04	UC2-NormColon
886	648782	1707.B12.gz43 216605	F	M00055942D:A01	UC2-NormColon
887	645900	1707.013.gz43_216634	F	M00055967B:F07	UC2-NormColon
888	648555	1707.D14.gz43_216639	F	M00055945C:D09	UC2-NormColon
889	452316	1707.J14.gz43_216645	F	M00055957B:F11	UC2-NormColon
890	461363	1707.D17.gz43_216687	F	M00055945D:D01	UC2-NormColon
891	644720	1707.D18.gz43_216703	F	M00055945D:E08	UC2-NormColon
892	647312	1707.L18.gz43_216711	F	M00055961D:F06	UC2-NormColon
893	646190	1707.N18.gz43_216713	F	M00055966B:H09	UC2-NormColon
894	531529	1707.H19.gz43_216723	F	M00055954C:G07	UC2-NormColon
895	393599	1707.E20.gz43_216736	F	M00055947C:E09	UC2-NormColon
896	645508	1707.N20.gz43_216745	F	M00055966C:D03	UC2-NormColon
897	383609	1707.O23.gz43_216794	F	M00055967D:C05	UC2-NormColon
898	447805	1707.B24.gz43_216797	F	M00055943B:E01	UC2-NormColon
899	694643	1707.P24.gz43_216811	F	M00055969A:F02	UC2-NormColon
900	647924	1708.I01.gz43_216820	F	M00055980B:B07	UC2-NormColon
901	645920	1708.P01.gz43_216827	F	M00055990A:F07	UC2-NormColon
902	641484	1708.M02.gz43_216840	F	M00055985D:E09	UC2-NormColon
903	640672	1708.A04.gz43_216860	F	M00055969B:B05	UC2-NormColon
904	644226	1708.K04.gz43_216870	F	M00055982C:H01	UC2-NormColon
905	557852	1708.C06.gz43_216894	F	M00055972C:C09	UC2-NormColon
906	558052	1708.C07.gz43_216910	F	M00055972C:F02	UC2-NormColon
907	644354	1708.I08.gz43_216932	F	M00055980C:G12	UC2-NormColon
908	645746	1708.L08.gz43_216935	F	M00055984D:E04	UC2-NormColon
909	647539	1708.M09.gz43_216952	F	M00055986A:F05	UC2-NormColon
910	645288	1708.L10.gz43_216967	F	M00055985A:B06	UC2-NormColon
911	645289	1708.A11.gz43_216972	F	M00055969D:B08	UC2-NormColon
912	642058	1708.G11.gz43_216978	F_	M00055978A:H03	UC2-NormColon
913	642955	1708.A12.gz43_216988	F _	M00055969D:F08	UC2-NormColon
914	640092	1708.H12.gz43_216995	F	M00055979B:G07	UC2-NormColon
915	558452	1708.P12.gz43_217003	F	M00055990D:B02	UC2-NormColon
916	645427	1708.L13.gz43_217015	F	M00055985B:C02	UC2-NormColon
917	649082	1708.B14.gz43_217021	F	M00055971B:A11	UC2-NormColon
918	451709	1708.K14.gz43_217030	F	M00055983C:C09	UC2-NormColon
919	640204	1708.H15.gz43_217043	F	M00055979C:B07	UC2-NormColon
920	640179	1708.N15.gz43_217049	F	M00055988A:A12	UC2-NormColon
921	582549	1708.P15.gz43_217051	F	M00055990D:G09	UC2-NormColon
922	484355	1708.G16.gz43_217058	F	M00055978B:F01	UC2-NormColon
923	646013	1708.J17.gz43_217077	F	M00055982A:G06	UC2-NormColon
924	649309	1708.P18.gz43_217099	F	M00055991A:D10	UC2-NormColon

Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
925	647333	1708.F19.gz43 217105	F	M00055977A:G09	UC2-NormColon
926	647058	1708,A21,gz43 217132	F	M00055970C;A05	UC2-NormColon
927	643723	1708.F21.gz43 217137	F	M00055977B:F06	UC2-NormColon
928	639359	1708.N21.gz43_217145	F	M00055988A:E05	UC2-NormColon
929	647539	1708.H23.gz43 217171	F	M00055980A:H06	UC2-NormColon
930	647211	1708.C24.gz43 217182	F	M00055973C:F10	UC2-NormColon
931	650018	1708.J24.gz43_217189	F	M00055982C:A12	UC2-NormColon
932	645848	1708.K24.gz43_217190	F	M00055984A:F05	UC2-NormColon
933	646212	1708.P24.gz43_217195	F	M00055991C:H11	UC2-NormColon
934	406436	1588.N01.gz43_217262	F	M00055553A:D08	UC2-NormColon
935	553651	1588,E02.gz43_217269	F	M00055542C:H05	UC2-NormColon
936	466092	1588.H02.gz43 217272	F	M00055546B:H03	UC2-NormColon
937	645891	1588.N02.gz43 217278	F	M00055553A:F11	UC2-NormColon
938	649390	1588.A03.gz43_217281	F	M00055536B:H11	UC2-NormColon
939	448358	1588.H04.gz43 217304	F	M00055546C:B01	UC2-NormColon
940	650982	1588.I04.gz43_217305	F	M00055547C:B07	UC2-NormColon
941	644012	1588.O05.gz43_217327	F	M00055554C:B04	UC2-NormColon
942	649099	1588.P06.gz43 217344	F	M00055556A:E07	UC2-NormColon
943	551662	1588.M07.gz43 217357	F	M00055551C:C08	UC2-NormColon
944	639316	1588.K08.gz43_217371	F	M00055549C:E10	UC2-NormColon
945	639480	1588.J09.gz43 217386	F	M00055548B:G06	UC2-NormColon
946	1318	1588.A11.gz43_217409	F	M00055537C:A01	UC2-NormColon
947	457146	1588.L11.gz43_217420	F	M00055550D:D02	UC2-NormColon
948	451671	1588.A15.gz43_217473	F	M00055537C:E04	UC2-NormColon
949	641401	1588,I15.gz43_217481	F	M00055547D:G06	UC2-NormColon
950	453606	1588.D17.gz43_217508	F	M00055542B:B11	UC2-NormColon
951	642414	1588.G17.gz43_217511	F	. M00055545D:C03	UC2-NormColon
952	649564	1588.H18.gz43_217528	F	M00055547A:H11	UC2-NormColon
953	529799	1588,I18.gz43_217529	F	M00055548A:D07	UC2-NormColon
954	645781	1588.N18.gz43_217534	F	M00055553D:E06	UC2-NormColon
955	447379	1588.M20.gz43_217565	F	M00055552C:G03	UC2-NormColon
956	639461	1588.A21.gz43_217569	F	M00055538B:G09	UC2-NormColon
957	451361	1588.D21.gz43_217572	F	M00055542C:B02	UC2-NormColon
958	449891	1588,I22.gz43_217593	F	M00055548A:F04	UC2-NormColon
959	639395	1588.O22.gz43_217599	F	M00055555D:B05	UC2-NormColon
960	644173	1588.F23.gz43_217606	F	M00055545A:C01	UC2-NormColon
961	648494	1588.H24.gz43_217624	F	M00055547B:G09	UC2-NormColon
962	477 0 64	1588.P24.gz43_217632	F	M00055557B:B10	UC2-NormColon
963	535436	1729.L02.gz43_217660	F	M00056092B:D10	UC2-NormColon
'964	650900	1729.B04.gz43_217682	F	M00056078B:G02	UC2-NormColon
965	649149	1729.D04.gz43_217684	F	M00056081A:C05	UC2-NormColon
966	644054	1729.K04.gz43_217691	F	M00056091A:E04	UC2-NormColon

Table 2

T	able 2				
SEQ			ÓRIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
967	649082	1729.P04.gz43_217696	F	M00056097B:C04	UC2-NormColon
968	454176	1729.J05.gz43_217706	F	M00056089D:A09	UC2-NormColon
969	448485	1729.A06.gz43_217713	F	M00056077B:E03	UC2-NormColon
970	643440	1729.N06.gz43_217726	F	M00056095B:A07	UC2-NormColon
971	649356	1729.J08.gz43 217754	F	M00056089D:E07	UC2-NormColon
972	710155	1729.L08.gz43_217756	F	M00056092C:E12	UC2-NormColon
973	560054	1729.E11.gz43_217797	F	M00056082A:E12	UC2-NormColon
974	644712	1729.K11.gz43_217803	F	M00056091C:C06	UC2-NormColon
975	646696	1729.A12.gz43_217809	F	M00056077C:H02	UC2-NormColon
976	644715	1729.H12.gz43_217816	F	M00056087B:G12	UC2-NormColon
977	644965	1729.N12.gz43_217822	F	M00056095C:E02	UC2-NormColon
978	645050	1729.K13.gz43_217835	F	M00056091C:D09	UC2-NormColon
979	649514	1729.N14.gz43_217854	F	M00056095C:G07	UC2-NormColon
980	570939	1729.E15.gz43_217861	F	M00056082B:G07	UC2-NormColon
981	647437	1729.F16.gz43_217878	F	M00056083D:D11	UC2-NormColon
982	650547	1729.L16.gz43_217884	F	M00056093A:B12	UC2-NormColon
983	649054	1729.P16.gz43_217888	F	M00056098B:C04	UC2-NormColon
984	645409	1729.K17.gz43_217899	F	M00056091D:C12	UC2-NormColon
985	613722	1729.B18.gz43_217906	F	M00056079A:B01	UC2-NormColon
986	418763	1729.L18.gz43_217916	F	M00056093A:F08	UC2-NormColon
987	619635	1729.K20.gz43_217947	F	M00056091D:E10	UC2-NormColon
988	645131	1729.L20.gz43_217948	F	M00056093B:D03	UC2-NormColon
989	451899	1729.M22.gz43_217981	F	M00056095A:C02	UC2-NormColon
990	597647	1729.P22.gz43_217984	F	M00056098D:A08	UC2-NormColon
991	560860	1717.F01.gz43_218406	F	M00055998B:D06	UC2-NormColon
992	640709	1717.P01.gz43_218416	F	M00056009D:C12	UC2-NormColon
993	447802	1717.O02.gz43_218431	F	M00056008B:G05	UC2-NormColon
994	641736	1717.I03.gz43_218441	F	M00056001D:B06	UC2-NormColon
995	464498	1717.J03.gz43_218442	F	M00056003B:G11	UC2-NormColon
996	452775	1717.K03.gz43_218443	F	M00056004D:F03	UC2-NormColon
997	561836	1717.L03.gz43_218444	F	M00056005C:F11	UC2-NormColon
998	444454	1717.K04.gz43_218459	F	M00056004D:F12	UC2-NormColon
999	639178	1717.O04.gz43_218463	F	M00056008C:D04	UC2-NormColon
1000	513888	1717.F05.gz43_218470	F	M00055998C:D04	UC2-NormColon
1001	643971	1717.I05.gz43_218473	F	M00056001D:G12	UC2-NormColon
1002	644824	1717.J05.gz43_218474	F	M00056003C;C09	UC2-NormColon
1003	448418	1717.N05.gz43_218478	F	M00056007C:A03	UC2-NormColon
1004	644030	1717.005.gz43_218479	F	M00056008C:D08	UC2-NormColon
1005	644972	1717.P05.gz43_218480	F	M00056010A:E05	UC2-NormColon
1006	649556	1717.B06.gz43_218482	F	M00055993D:F12	UC2-NormColon
1007	644611	1717.I06.gz43_218489	F	M00056002A:A03	UC2-NormColon
1008	645162	1717.J07.gz43_218506	F	M00056003C:H10	UC2-NormColon

Т	able 2				
SEQ			ORIEN		
	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
1009	470462	1717.L07.gz43_218508	F	M00056005D:C04	UC2-NormColon
1010	645662	1717.A08.gz43_218513	F	M00055992A:D08	UC2-NormColon
1011	649259 650018	1717.H08.gz43_218520 1717.D09.gz43 218532	F	M00056000C:D09	UC2-NormColon
1012	640645	1717.D09.gz43_218532 1717.D10.gz43_218548	F	M00055996C:A02 M00055996C:A08	UC2-NormColon UC2-NormColon
1013	643477	1717.M10.gz43_218557	F	M00055996C:A08	UC2-NormColon
1015	541214	1717.E11.gz43 218565	F	M00055997D:A11	UC2-NormColon
1016	651083	1717.M11.gz43 218573	F	M00056006D:F05	UC2-NormColon
1017	549889	1717.G12.gz43_218583	F	M00055999D:C05	UC2-NormColon
1018	558856	1717.D13.gz43 218596	F	M00055996D:D07	UC2-NormColon
1019	644022	1717.N13.gz43_218606	F	M00056007D:F07	UC2-NormColon
1020	644612	1717.O13.gz43 218607	F	M00056008D:D02	UC2-NormColon
1021	644949	1717.P13.gz43 218608	F	M00056011A:C11	UC2-NormColon
1022	643808	1717.C14.gz43_218611	F	M00055995A:G09	UC2-NormColon
1023	642260	1717.O14.gz43_218623	F	M00056008D:D03	UC2-NormColon
1024	651119	1717.B15.gz43_218626	F	M00055994B:B12	UC2-NormColon
1025	477521	1717.C15.gz43_218627	F	M00055995B:A02	UC2-NormColon
1026	587106	1717.D15.gz43_218628	F	M00055996D:G02	UC2-NormColon
1027	419443	1717.I15.gz43_218633	F	M00056002D:E09	UC2-NormColon
1028	643440	1717.L15.gz43_218636	F	M00056006A:E04	UC2-NormColon
1029	555736	1717.M16.gz43_218653	F	M00056007A:B02	UC2-NormColon
1030	644210	1717.G17.gz43_218663	F	M00055999D:H04	UC2-NormColon
1031	449261	1717.I17.gz43_218665	F	M00056003A:A08	UC2-NormColon
1032	643498	1717.J17.gz43_218666	F	M00056004B:F04	UC2-NormColon
1033	517274	1717.P17.gz43_218672	F	M00056011B:A06	UC2-NormColon
1034	532307 643288	1717.L18.gz43_218684 1717.A19.gz43_218689	F	M00056006A:G09 M00055993A:E02	UC2-NormColon
1035	478025	1717.F19.gz43_218694	F	M00055999A:A04	UC2-NormColon UC2-NormColon
1037	447822	1717.G19.gz43_218695	F	M00056000A:B02	UC2-NormColon
1037	641580	1717.119.gz43_218697	F	M00056003A:B07	UC2-NormColon
1039	489426	1717.P19.gz43 218704	F	M00056011B:E10	UC2-NormColon
1040	650534	1717.C20.gz43 218707	F	M00055995C:F05	UC2-NormColon
1041	644990	1717.E20.gz43 218709	F	M00055998A;F06	UC2-NormColon
1042	207530	1717.F20.gz43 218710	F	M00055999A:B07	UC2-NormColon
1043	645197	1717.I20.gz43 218713	F	M00056003A:C01	UC2-NormColon
1044	414739	1717.K20.gz43 218715	F	M00056005B:H08	UC2-NormColon
1045	592941	1717.L20.gz43_218716	F	M00056006B:B05	UC2-NormColon
1046	447556	1717.H21.gz43_218728	F	M00056001C:F07	UC2-NormColon
1047	238586	1717.N21.gz43_218734	F	M00056008B:E04	UC2-NormColon
1048	644612	1717.E22.gz43_218741	F	M00055998B:A04	UC2-NormColon
1049	646387	1717.F22.gz43 218742	F	M00055999A:H10	UC2-NormColon
1050	647443	1717.G22.gz43_218743	F	M00056000A:E11	UC2-NormColon

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Table 2

1	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	Т	CLONE ID	LIBRARY
1051	555172	1717.P22.gz43 218752	F	M00056011C:D04	UC2-NormColon
1052	552846	1717.D23.gz43_218756	F	M00055997A:F04	UC2-NormColon
1053	639377	1717.B24.gz43 218770	F	M00055994C:F05	UC2-NormColon
1054	449713	1717.M24.gz43_218781	F	M00056007B:C10	UC2-NormColon
1055	456627	1717.N24.gz43_218782	F	M00056008B:G01	UC2-NormColon
1056	646143	1717.O24.gz43_218783	F	M00056009D:A02	UC2-NormColon
1057	639629	1718.H03.gz43_218827	F	M00056021C:H08	UC2-NormColon
1058	643744	1718.D04.gz43_218839	F	M00056016C:F11	UC2-NormColon
1059	397399	1718.P04.gz43_218851	F	M00056032B:H08	UC2-NormColon
1060	650391	1718.G06.gz43_218874	F	M00056020C:H03	UC2-NormColon
1061	447659	1718.N06.gz43 218881	F	M00056029D:D08	UC2-NormColon
1062	644967	1718.D07.gz43 218887	F	M00056016D:E02	UC2-NormColon
1063	545509	1718.G08.gz43 218906	F	M00056020D:D03	UC2-NormColon
1064	643975	1718.D09.gz43 218919	F	M00056017A:B08	UC2-NormColon
1065	645883	1718.M11.gz43 218960	F	M00056028D:F06	UC2-NormColon
1066	538808	1718.B12.gz43 218965	F	M00056013A:C09	UC2-NormColon
1067	425455	1718.P13.gz43 218995	F	M00056033A:C11	UC2-NormColon
1068	467057	1718.M20.gz43 219104	F	M00056029B:H03	UC2-NormColon
1069	640181	1718.B21.gz43 219109	F	M00056013D:D07	UC2-NormColon
1070	650348	1718.F23.gz43_219145	F	M00056020B:C09	UC2-NormColon
1071	451993	1718.P24.gz43_219171	F	M00056033C:H09	UC2-NormColon
1072	647976	1719.E01.gz43_219176	F	M00056039A:F04	UC2-NormColon
1073	557676	1719.I01.gz43 219180	F	M00056045A:D03	UC2-NormColon
1074	644639	1719.N01.gz43_219185	F	M00056050C:A09	UC2-NormColon
1075	644738	1719.J02.gz43_219197	F	M00056046A:B04	UC2-NormColon
1076	645973	1719.A03.gz43_219204	F	M00056033D:F06	UC2-NormColon
1077	550376	1719.B03.gz43_219205	F	M00056035B:A04	UC2-NormColon
1078	646088	1719.D03.gz43_219207	F	M00056037B:G02	UC2-NormColon
1079	506920	1719.P03.gz43_219219	F	M00056053A:F01	UC2-NormColon
1080	642198	1719.A04.gz43_219220	F	M00056033D:G07	UC2-NormColon
1081	649293	1719.E04.gz43_219224	F	M00056039B:C03	UC2-NormColon
1082	641193	1719.H04.gz43_219227	F	M00056043D;E03	UC2-NormColon
1083	461517	1719.J04.gz43_219229	F	M00056046A:B12	UC2-NormColon
1084	645070	1719.K04.gz43_219230	F	M00056046D:C11	UC2-NormColon
1085	645942	1719.M04.gz43_219232	F	M00056049B;E11	UC2-NormColon
1086	645151	1719.O05.gz43_219250	F	M00056051D:H02	UC2-NormColon
1087	566745	1719.B06.gz43_219253	F	M00056035B:D11	UC2-NormColon
1088	642184	1719.M06.gz43_219264	F	M00056049C:A08	UC2-NormColon
1089	649429	1719.O06.gz43_219266	F	M00056052A;A11	UC2-NormColon
1090	549786	1719.B07.gz43_219269	F	M00056035B:E10	UC2-NormColon
1091	452102	1719.J07.gz43_219277	F	M00056046A:G08	UC2-NormColon
1092	644819	1719.O07.gz43 219282	F	M00056052A;C07	UC2-NormColon

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	able 2				Г
SEQ			ORIEN		
	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
1093	453804	1719.E08.gz43 219288	F	M00056039C:D05	UC2-NormColon
1094	650348	1719,K09,gz43 219310	F	M00056047A:E09	UC2-NormColon
1095	644894	1719.L09.gz43 219311	F	M00056048A:D12	UC2-NormColon
1096	646048	1719.A10.gz43 219316	F	M00056034A:G10	UC2-NormColon
1097	638912	1719,D10,gz43 219319	F	M00056037C:G11	UC2-NormColon
1098	644817	1719.F10.gz43 219321	F	M00056041B;F03	UC2-NormColon
1099	645973	1719.E11.gz43_219336	F	M00056039C:G05	UC2-NormColon
1100	643748	1719.M11.gz43 219344	F	M00056049C;H04	UC2-NormColon
1101	649259	1719.N11.gz43 219345	F	M00056051A:C03	UC2-NormColon
1102	642852	1719.E12.gz43 219352	F	M00056039C:H01	UC2-NormColon
1103	451544	1719.L12.gz43 219359	F	M00056048B:E01	UC2-NormColon
1104	528616	1719.N12.gz43 219361	F	M00056051A:H11	UC2-NormColon
1105	460727	1719.G13.gz43 219370	F	M00056043A:A09	UC2-NormColon
1105	468689	1719.C14.gz43_219370	F	M00056036D;B06	UC2-NormColon
1107	452729	1719.C14.gz43_219382 1719.C15.gz43_219398	F		
1107	481614	1719.C13.gz43_219398 1719.D15.gz43_219399	F	M00056036D:D01	UC2-NormColon
			F	M00056038B:G08	UC2-NormColon
1109	649900	1719.H15.gz43_219403		M00056044C:C01	UC2-NormColon
1110	645375	1719.N15.gz43_219409	F	M00056051B:B03	UC2-NormColon
1111	497101	1719.P15.gz43_219411	F	M00056053D:D07	UC2-NormColon
1112	556325	1719.G16.gz43_219418	F	M00056043A:H12	UC2-NormColon
1113	645431	1719.M16.gz43_219424	F	M00056049D:D03	UC2-NormColon
1114	452325	1719.A17.gz43_219428	F	M00056034C:D07	UC2-NormColon
1115	447645	1719.B17.gz43_219429	F	M00056035D:C08	UC2-NormColon
1116	644054	1719.C17.gz43_219430	F	M00056036D:F05	UC2-NormColon
1117	641715	1719.M17.gz43_219440	F	M00056049D:E01	UC2-NormColon
1118	644692	1719.P17.gz43_219443	F	M00056054A:B06	UC2-NormColon
1119	465589	1719.P18.gz43_219459	F	M00056054A:D09	UC2-NormColon
1120	643897	1719.G19.gz43_219466	F	M00056043B:E03	UC2-NormColon
1121	456840	1719.L19.gz43_219471	F	M00056048D:B09	UC2-NormColon
1122	467255	1719.P19.gz43 219475	F	M00056054A:E03	UC2-NormColon
1123	645530	1719.C20.gz43_219478	F	M00056037A:D11	UC2-NormColon
1124	648390	1719.A21.gz43_219492	F	M00056034C:H09	UC2-NormColon
1125	467597	1719.B21.gz43 219493	F	M00056035D:G10	UC2-NormColon
1126	468330	1719,D21.gz43 219495	F	M00056038D:F03	UC2-NormColon
1127	454438	1719.E21.gz43 219496	F	M00056040C:B03	UC2-NormColon
1128	402476	1719.I21.gz43 219500	F	M00056045D:C09	UC2-NormColon
1129	558439	1719.M21.gz43 219504	F	M00056050A;D02	UC2-NormColon
1130	595066	1719.N21.gz43 219505	F	M00056051C;C09	UC2-NormColon
1131	646687	1719.E22.gz43 219512	F	M00056040C:C12	UC2-NormColon
1132	452026	1719.H22.gz43 219515	F	M00056044D:F01	UC2-NormColon
1133	639255	1719.K22.gz43 219518	F	M00056047C:E03	UC2-NormColon
1134	612572	1719.L22.gz43 219519	F	M00056049A;C11	UC2-NormColon

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T	able 2				
SEO			ORIEN		
	CLUSTER	SEO NAME	T	CLONE ID	LIBRARY
1135	641786	1719.O22.gz43 219522	F	M00056052D:G01	UC2-NormColon
1136	647306	1719.P22.gz43 219523	F	M00056054A:G07	UC2-NormColon
1137	640559	1719.A23.gz43 219524	F	M00056034D:E09	UC2-NormColon
1138	550365	1719.D23.gz43 219527	F	M00056039A:A02	UC2-NormColon
1139	462293	1719.I23.gz43 219532	F	M00056045D:E06	UC2-NormColon
1140	587106	1719.N23.gz43 219537	F	M00056051C:H09	UC2-NormColon
1141	483266	1719.F24.gz43 219545	F	M00056042B:B05	UC2-NormColon
1142	490898	1719.G24.gz43 219546	F	M00056043C:G03	UC2-NormColon
1143	640603	1719.N24.gz43 219553	F	M00056051D:A07	UC2-NormColon
1144	643510	1720.G01.gz43 219562	F	M00056062D:F12	UC2-NormColon
1145	645049	1720.A02.gz43 219572	F	M00056054B:G05	UC2-NormColon
1146	466795	1720,C02,gz43 219574	F	M00056057B:D01	UC2-NormColon
1147	14157	1720.I02.gz43 219580	F	M00056066B:H04	UC2-NormColon
1148	558494	1720.A04.gz43 219604	F	M00056054C:C09	UC2-NormColon
1149	644927	1720.C04.gz43 219606	F	M00056057B:E12	UC2-NormColon
1150	648688	1720.C05.gz43 219622	F	M00056057C:B02	UC2-NormColon
1151	454906	1720.A06.gz43 219636	F	M00056054C:E12	UC2-NormColon
1152	644597	1720.D06.gz43 219639	F	M00056059A:A02	UC2-NormColon
1153	460047	1720.K06.gz43 219646	F	M00056069A:F01	UC2-NormColon
1154	62053	1720.C08.gz43 219670	F	M00056057C:D06	UC2-NormColon
1155	649054	1720.D08.gz43 219671	F	M00056059A:B01	UC2-NormColon
1156	644225	1720.K08.gz43 219678	F	M00056069A:F06	UC2-NormColon
1157	447635	1720.C09.gz43 219686	F	M00056057C:E01	UC2-NormColon
1158	642166	1720.E10.gz43 219704	F	M00056060D:H11	UC2-NormColon
1159	473007	1720.D11.gz43 219719	F	M00056059B:D10	UC2-NormColon
1160	551181	1720,M11,gz43 219728	F	M00056072B:C06	UC2-NormColon
1161	644928	1720.C12.gz43 219734	F	M00056057C:E12	UC2-NormColon
1162	644916	1720.E12.gz43_219736	F	M00056061A:E05	UC2-NormColon
1163	644701	1720.L12.gz43_219743	F	M00056070C:B09	UC2-NormColon
1164	487567	1720.O12.gz43_219746	F	M00056074C:H08	UC2-NormColon
1165	641801	1720.E13.gz43 219752	F	M00056061B:A03	UC2-NormColon
1166	651108	1720.O13.gz43_219762	F	M00056074C:H09	UC2-NormColon
1167	447676	1720.A14.gz43_219764	F	M00056055A:E04	UC2-NormColon
1168	645076	1720.K14.gz43 219774	F	M00056069B:E04	UC2-NormColon
1169	641469	1720.C15.gz43_219782	F	M00056057D:E11	UC2-NormColon
1170	375577	1720.G15.gz43_219786	F	M00056063C:F10	UC2-NormColon
1171	558154	1720.C16.gz43_219798	F	M00056058A:B07	UC2-NormColon
1172	650549	1720.G16.gz43_219802	F	M00056063C:G11	UC2-NormColon
1173	461351	1720.I16.gz43_219804	F	M00056066D:G04	UC2-NormColon
1174	640818	1720.A18.gz43_219828	F	M00056055B:B06	UC2-NormColon
1175	455716	1720.I18.gz43_219836	F	M00056067A:A12	UC2-NormColon
1176	644822	1720,J18.gz43_219837	F	M00056068D:A06	UC2-NormColon

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1	able 2				
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ID NO	CLUSTER	SEQ NAME	Т	CLONE ID	LIBRARY
1177	467780	1720.C19.gz43 219846	F	M00056058A:F08	UC2-NormColon
1178	446415	1720.F20.gz43_219865	F	M00056062D:D08	UC2-NormColon
1179	553923	1720.H20.gz43_219867	F	M00056066A:D03	UC2-NormColon
1180	645900	1720.I21.gz43 219884	F	M00056067B:D08	UC2-NormColon
1181	649364	1720.M21.gz43_219888	F	M00056072D:E08	UC2-NormColon
1182	643348	1720.N23.gz43_219921	F	M00056074B:D12	UC2-NormColon
1183	648654	1731.A01.gz43_219940	F	M00056120C:H04	UC2-NormColon
1184	648547	1731.C01.gz43_219942	F	M00056123B:G05	UC2-NormColon
1185	463824	1731.I01.gz43_219948	F	M00056133D:D09	UC2-NormColon
1186	648721	1731.N01.gz43_219953	F	M00056140C:E04	UC2-NormColon
1187	648481	1731.001.gz43_219954	F	M00056142A:F03	UC2-NormColon
1188	644037	1731.A02.gz43_219956	F	M00056120D:F01	UC2-NormColon
1189	635354	1731.H02.gz43_219963	F	M00056132B;C12	UC2-NormColon
1190	711493	1731.J02.gz43 219965	F	M00056134D:G07	UC2-NormColon
1191	648742	1731.L02.gz43_219967	F	M00056137D;H05	UC2-NormColon
1192	476380	1731.F03.gz43_219977	F	M00056129C:F09	UC2-NormColon
1193	649617	1731.J03.gz43_219981	F	M00056134D:G11	UC2-NormColon
1194	639555	1731.B04.gz43_219989	F	M00056122B:A07	UC2-NormColon
1195	465127	1731.D04.gz43_219991	F	M00056125B:F01	UC2-NormColon
1196	458938	1731.L04.gz43_219999	F	M00056138A:B11	UC2-NormColon
1197	453726	1731.G05.gz43_220010	F	M00056131A:E09	UC2-NormColon
1198	557026	1731.H06.gz43_220027	F	M00056132C:F04	UC2-NormColon
1199	422242	1731.D07.gz43_220039	F	M00056127A:A10	UC2-NormColon
1200	456985	1731.E07.gz43_220040	F	M00056128B:A07	UC2-NormColon
1201	640695	1731.I07.gz43_220044	F	M00056134A:D04	UC2-NormColon
1202	627515	1731.B08.gz43_220053	F	M00056122B:G09	UC2-NormColon
1203	648341	1731.M08.gz43_220064	F	M00056139D:E05	UC2-NormColon
1204	464990	1731_A09.gz43_220068	F	M00056121A:E05	UC2-NormColon
1205	459521	1731.J10.gz43_220093	F	M00056135B:E03	UC2-NormColon
1206	549691	1731.L10.gz43_220095	F	M00056138B:F05	UC2-NormColon
1207	514838	1731.M10.gz43_220096	F	M00056139D:H04	UC2-NormColon
1208	452738	1731.O10.gz43_220098	F	M00056142C:A09	UC2-NormColon
1209	644928	1731.D11.gz43_220103	F	M00056127A:H03	UC2-NormColon
1210	460190	1731.B12.gz43_220117	F	M00056122C:H12	UC2-NormColon
1211	562378	1731.G12.gz43_220122	F	M00056131B:E01	UC2-NormColon
1212	647949	1731.P12.gz43_220131	F	M00056144D:C05	UC2-NormColon
1213	557419	1731.B13.gz43_220133	F	M00056122D:B07	UC2-NormColon
1214	638934	1731.D13.gz43_220135	F	M00056127B:E11	UC2-NormColon
1215	646785	1731.F13.gz43_220137	F	M00056130B:E11	UC2-NormColon
1216	647248	1731.E14.gz43_220152	F	M00056128C:F02	UC2-NormColon
1217	470006	1731.I14.gz43_220156	F	M00056134C:C06	UC2-NormColon
1218	648311	1731.K14.gz43_220158	F	M00056137B:F07	UC2-NormColon

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SEQ			ORIEN		
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1219	649299	1731.F15.gz43_220169	F	M00056130C:C12	UC2-NormColon
1220	478393	1731.K15.gz43_220174	F	M00056137B:F11	UC2-NormColon
1221	647290	1731.F16.gz43_220185	F	M00056130C:F08	UC2-NormColon
1222	649735	1731.G16.gz43_220186	F	M00056131C:H03	UC2-NormColon
1223	646039	1731.J16.gz43_220189	F	M00056135C:C05	UC2-NormColon
1224	646581	1731.M16.gz43_220192	F	M00056140A:F12	UC2-NormColon
1225	470667	1731.N16.gz43_220193	_ F	M00056141C:H01	UC2-NormColon
1226	642650	1731.J17.gz43_220205	F	M00056135C:C08	UC2-NormColon
1227	642275	1731.K17.gz43_220206	F	M00056137C:A01	UC2-NormColon
1228	651015	1731.L17.gz43_220207	F	M00056138D:G08	UC2-NormColon
1229	642936	1731.H18.gz43_220219	F	M00056133B:F01	UC2-NormColon
1230	642204	1731.A19.gz43_220228	F	M00056121D:A12	UC2-NormColon
1231	557867	1731.J19.gz43_220237	F	M00056135C:H06	UC2-NormColon
1232	639395	1731.K19.gz43_220238	F	M00056137C:E01	UC2-NormColon
1233	645367	1731.K20.gz43_220254	F	M00056137C:G02	UC2-NormColon
1234	647232	1731.L20.gz43_220255	F	M00056139A:D12	UC2-NormColon
1235	647431	1731.E21.gz43_220264	F	M00056129A:G01	UC2-NormColon
1236	708175	1731.I21.gz43_220268	F	M00056134D:B07	UC2-NormColon
1237	503122	1731.A22.gz43_220276	F	M00056122A:A05	UC2-NormColon
1238	515350	1731.C22.gz43_220278	F	M00056124D:F06	UC2-NormColon
1239	556336	1731.F22.gz43_220281	F	M00056130D:E06	UC2-NormColon
1240	589098	1731.A23.gz43_220292	F	M00056122A:B05	UC2-NormColon
1241	531461	1731.B23.gz43_220293	F	M00056123B:F02	UC2-NormColon
1242	455808	1731.C23.gz43_220294	F	M00056124D:G03	UC2-NormColon
1243	643510	1731,D23.gz43_220295	F	M00056127D:G10	UC2-NormColon
1244	648996	1731.J23.gz43_220301	F	M00056136A:B11	UC2-NormColon
1245	446164	1731.N23.gz43_220305	F	M00056141D:E08	UC2-NormColon
1246	419114	1731.O23.gz43_220306 .	F	M00056143A:H08	UC2-NormColon
1247	648076	1731.A24.gz43_220308	F	M00056122A:D02	UC2-NormColon
1248	649482	1731.I24.gz43_220316	F	M00056134D:F02	UC2-NormColon
1249	646323	1732.H01.gz43_220331	F	M00056155B:A02	UC2-NormColon
1250	550714	1732.A05.gz43_220388	F	M00056146A:A09	UC2-NormColon
1251	649482	1732.J05.gz43_220397	F	M00056158C:D11	UC2-NormColon
1252	650987	1732.J06.gz43_220413	F	M00056158C:F09	UC2-NormColon
1253	644221	1732.F07.gz43_220425	F	M00056153A;G04	UC2-NormColon
1254	647412	1732.M07.gz43_220432	F	M00056162C:F02	UC2-NormColon
1255	416624	1732.I08.gz43_220444	F	M00056157A:B11	UC2-NormColon
1256	644801	1732.F09.gz43_220457	F	M00056153A:H07	UC2-NormColon
1257	650249	1732.G10.gz43_220474	F	M00056154B:F11	UC2-NormColon
1258	465734	1732.C11.gz43_220486	F	M00056149C:A02	UC2-NormColon
1259	605761	1732.E11.gz43_220488	F	M00056152A:B11	UC2-NormColon
1260	649782	1732.M11.gz43_220496	F	M00056162D:A01	UC2-NormColon

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Table 2

	able 2				
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ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
1261	449956	1732.A12.gz43 220500	F	M00056146A:H09	UC2-NormColon
1262	642244	1732.I12.gz43 220508	F	M00056157B:A08	UC2-NormColon
1263	650348	1732,L12.gz43 220511	F	M00056161C:G06	UC2-NormColon
1264	461135	1732.E13.gz43_220520	F	M00056152A:D07	UC2-NormColon
1265	16092	1732.L13.gz43 220527	F	M00056161C:H10	UC2-NormColon
1266	650116	1732.M13.gz43_220528	F	M00056162D:D03	UC2-NormColon
1267	650534	1732.O13.gz43_220530	F	M00056165C:B11	UC2-NormColon
1268	640634	1732.A14.gz43_220532	F	M00056146B:E05	UC2-NormColon
1269	446184	1732,I14.gz43_220540	F	M00056157B:H06	UC2-NormColon
1270	380477	1732.M14.gz43 220544	F	M00056162D:D06	UC2-NormColon
1271	463966	1732.K15.gz43_220558	F	M00056160C:F12	UC2-NormColon
1272	649578	1732.O15.gz43_220562	F	M00056165D:D09	UC2-NormColon
1273	650877	1732.C16.gz43_220566	F	M00056149C:E10	UC2-NormColon
1274	449882	1732.F16.gz43_220569	F	M00056153B:F11	UC2-NormColon
1275	650909	1732.J16.gz43_220573	F	M00056159A:E03	UC2-NormColon
1276	645100	1732.K16.gz43_220574	F	M00056160C:G01	UC2-NormColon
1277	649883	1732.M17.gz43_220592	F	M00056163A:B10	UC2-NormColon
1278	554581	1732.I19.gz43_220620	F	M00056157C:E10	UC2-NormColon
1279	467972	1732.L19.gz43_220623	F	M00056162A:B06	UC2-NormColon
1280	488592	1732.A20.gz43_220628	F	M00056146D:D04	UC2-NormColon
1281	651075	1732.E20.gz43_220632	F	M00056152C:G08	UC2-NormColon
1282	649991	1732.K20.gz43_220638	F	M00056160D:C06	UC2-NormColon
1283	649947	1732,L20.gz43_220639	F	M00056162A:C09	UC2-NormColon
1284	224092	1732.N20.gz43_220641	F	M00056164A:H03	UC2-NormColon
1285	555634	1732.L21.gz43_220655	F	M00056162A:E09	UC2-NormColon
1286	639485	1732.C22.gz43_220662	F	M00056149D:F06	UC2-NormColon
1287	646596	1732.I23.gz43_220684	F	M00056157D:H08	UC2-NormColon
1288	459260	1732.J23.gz43_220685	F	M00056159C:C11	UC2-NormColon
1289	634409	1732.L23.gz43_220687	F	M00056162A:F01	UC2-NormColon
1290	650820	1732.C24.gz43_220694	F	M00056150A:E04	UC2-NormColon
1291	559349	1741.M01.gz43_220720	F	M00056184C:C07	UC2-NormColon
1292	650063	1741.D02.gz43_220727	F	M00056171C:A08	UC2-NormColon
1293	492982	1741.J02.gz43_220733	F	M00056180B:D05	UC2-NormColon
1294	557353	1741.K02.gz43_220734	F	M00056182A:B04	UC2-NormColon
1295	649299	1741.G03.gz43_220746	F	M00056175D:E05	UC2-NormColon
1296	648063	1741.F04.gz43_220761	F	M00056174C;H09	UC2-NormColon
1297	641193	1741.J04.gz43_220765	F	M00056180B:F11	UC2-NormColon
1298	561856	1741.L04.gz43_220767	F	M00056183B;G11	UC2-NormColon
1299	642631	1741.M04.gz43_220768	F	M00056184C:E08	UC2-NormColon
1300	648344	1741.P04.gz43_220771	F	M00056188B:E07	UC2-NormColon
1301	458085	1741.B05.gz43_220773	F	M00056169C:F07	UC2-NormColon
1302	394772	1741.C06.gz43_220790	F	M00056170C;C09	UC2-NormColon

Table 2

	able 2				
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1304	593173	1741.G07.gz43 220810	F	M00056176A:D06	UC2-NormColon
1305	643206	1741.K07.gz43 220814	F	M00056182A:G11	UC2-NormColon
1306	646309	1741,M07,gz43 220816	F	M00056184C:H03	UC2-NormColon
1307	553023	1741.N07.gz43 220817	F	M00056185D:C01	UC2-NormColon
1308	649705	1741.O07.gz43 220818	F	M00056186D:G05	UC2-NormColon
1309	555771	1741,B08.gz43 220821	F	M00056169D;A03	UC2-NormColon
1310	379805	1741.I08.gz43 220828	F	M00056179A:D12	UC2-NormColon
1311	710362	1741.L08.gz43 220831	F	M00056183C:F01	UC2-NormColon
1312	650188	1741.P08.gz43 220835	F	M00056188C:D02	UC2-NormColon
1313	649795	1741.A09.gz43 220836	F	M00056168C:A05	UC2-NormColon
1314	460506	1741.K09.gz43 220846	F	M00056182B:G12	UC2-NormColon
1315	502683	1741.N09.gz43 220849	F	M00056185D:D06	UC2-NormColon
1316	453768	1741.F10.gz43 220857	F	M00056175A:A06	UC2-NormColon
1317	457272	1741.G10.gz43 220858	F	M00056176B:B05	UC2-NormColon
1318	378694	1741.J10.gz43 220861	F	M00056180D:E06	UC2-NormColon
1319	649136	1741.F11.gz43 220873	F	M00056175A;C04	UC2-NormColon
1320	555655	1741.M11.gz43 220880	F	M00056184D:D06	UC2-NormColon
1321	648777	1741.B12.gz43 220885	F	M00056169D:D05	UC2-NormColon
1322	446289	1741.G12.gz43 220890	F	M00056176B:E10	UC2-NormColon
1323	449452	1741.J12.gz43_220893	F	M00056180D:F05	UC2-NormColon
1324	648665	1741.P12.gz43 220899	F	M00056188C:H04	UC2-NormColon
1325	561747	1741.D14.gz43_220919	F	M00056172A:H06	UC2-NormColon
1326	646915	1741.E14.gz43_220920	F	M00056174B;D02	UC2-NormColon
1327	648865	1741.G14.gz43_220922	F	M00056176C:B11	UC2-NormColon
1328	639886	1741.I14.gz43_220924	F	M00056179B:H12	UC2-NormColon
1329	183842	1741.J14.gz43_220925	F	M00056180D:G07	UC2-NormColon
1330	469409	1741.K14.gz43_220926	F	M00056182C:F08	UC2-NormColon
1331	649702	1741.L14.gz43_220927	F	M00056184A:B11	UC2-NormColon
1332	646581	1741.N14.gz43_220929	F	M00056186A:B09	UC2-NormColon
1333	638730	1741.H15.gz43_220939	_ F	M00056178B:A11	UC2-NormColon
1334	641525	1741.K15.gz43_220942	F	M00056182D:A07	UC2-NormColon
1335	650800	1741.B16.gz43_220949	F	M00056169D:H05	UC2-NormColon
1336	647086	1741.E16.gz43_220952	F	M00056174B:E02	UC2-NormColon
1337	471836	1741.J16.gz43_220957	F	M00056181A:B05	UC2-NormColon
1338	640285	1741.E17.gz43_220968	F	M00056174B:E04	UC2-NormColon
1339	459923	1741.H17.gz43_220971	F	M00056178B:H08	UC2-NormColon
1340	452729	1741.J17.gz43 220973	F	M00056181A:E07	UC2-NormColon
1341	209414	1741.H18.gz43_220987	F	M00056178C:C02	UC2-NormColon
1342	463487	1741.L18.gz43_220991	F	M00056184B:D08	UC2-NormColon
1343	645288	1741.F19.gz43_221001	F	M00056175C:A10	UC2-NormColon
1344	640514	1741.J19.gz43 221005	F	M00056181B:B02	UC2-NormColon

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Table 2

T	able 2				
SEQ	CV LIGHTED	(TO) / I / I	ORIEN		
	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
1345	459974	1741.K19.gz43_221006	F	M00056183A:B09	UC2-NormColon
1346	465594	1741.P19.gz43_221011	F	M00056189B:A01	UC2-NormColon
1347	646173	1741.C20.gz43_221014	F	M00056171A:H01	UC2-NormColon
1348	557010	1741.E20.gz43_221016	F	M00056174B:H11	UC2-NormColon
1349	650146	1741.F20.gz43_221017	F	M00056175C:B11	UC2-NormColon
1350	499690	1741.G20.gz43_221018	F	M00056177A:D12	UC2-NormColon
1351	584745	1741.H20.gz43_221019	F	M00056178C:E12	UC2-NormColon
1352	446987	1741.L20.gz43_221023	F	M00056184B:H08	UC2-NormColon
1353	458683	1741.A21.gz43_221028	F	M00056169B:C02	UC2-NormColon
1354	640922	1741.H21.gz43_221035	F	M00056178C;F02	UC2-NormColon
1355	561793	1741.J21.gz43_221037	F	M00056181B:E11	UC2-NormColon
1356	641945	1741.021.gz43_221042	F	M00056188A:E05	UC2-NormColon
1357	645578	1741.J22.gz43_221053	F	M00056181C:D02	UC2-NormColon
1358	647688	1741.L22.gz43_221055	F	M00056184C:A09	UC2-NormColon
1359	610986	1741.G23.gz43_221066	F	M00056177A:H01	UC2-NormColon
1360	644053	1741.I23.gz43_221068	F	M00056180B:A06	UC2-NormColon
1361	639132	1741.M23.gz43_221072	F	M00056185B:B06	UC2-NormColon
1362	642256	1741.B24.gz43_221077	F	M00056170B:B09	UC2-NormColon
1363	648820	1741.F24.gz43_221081	F	M00056175D:A06	UC2-NormColon
1364	650975	1741.L24.gz43_221087	F	M00056184C:C03	UC2-NormColon
1365	644801	1741.N24.gz43_221089	F	M00056186B:H09	UC2-NormColon
1366	637387	1742.G01.gz43_221098	F	M00056198A:B05	UC2-NormColon
1367	642118	1742.P01.gz43_221107	F	M00056212C:C09	UC2-NormColon
1368	397363	1742.F03.gz43_221129	F	M00056196C:B03	UC2-NormColon
1369	463168	1742.J03.gz43_221133	F	M00056203D:F05	UC2-NormColon
1370	172843	1742.M03.gz43_221136	F	M00056208B:C10	UC2-NormColon
1371	570939	1742.J04.gz43_221149	F	M00056203D:H03	UC2-NormColon
1372	463545	1742.L04.gz43_221151	F	M00056206D:F07	UC2-NormColon
1373	456920	1742.J05.gz43_221165	F	M00056204A:B08	UC2-NormColon
1374	644916	1742.E06.gz43_221176	F	M00056195C:F12	UC2-NormColon
1375	559955	1742.M06.gz43_221184	F	M00056208C:A10	UC2-NormColon
1376	453762	1742.B08.gz43_221205	F	M00056191B:E08	UC2-NormColon
1377	640913	1742.I08.gz43_221212	F	M00056201C:C03	UC2-NormColon
1378	648590	1742.F09.gz43 221225	F	M00056196C:G12	UC2-NormColon
1379	470801	1742.N09.gz43 221233	F	M00056210B:C11	UC2-NormColon
1380	645223	1742.M11.gz43_221264	F	M00056208D:A12	UC2-NormColon
1381	649402	1742.E12.gz43 221272	F	M00056195D:D07	UC2-NormColon
1382	649354	1742.N12.gz43 221281	F	M00056210B:E03	UC2-NormColon
1383	650756	1742.K15.gz43_221326	F	M00056205D:G04	UC2-NormColon
1384	648754	1742.J18.gz43 221373	F	M00056204C:H12	UC2-NormColon
1385	645177	1742.K19.gz43 221390	F	M00056206B:A10	UC2-NormColon
1386	507050	1742.M19.gz43 221392	F	M00056209B:D02	UC2-NormColon

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T	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
1387	646275	1742.P19.gz43_221395	F	M00056213C:C01	UC2-NormColon
1388	574166	1743.H01.gz43_221483	F	M00056223A:D10	UC2-NormColon
1389	649709	1743.H03.gz43_221515	F	M00056223A:H07	UC2-NormColon
1390	648155	1743.C04.gz43_221526	F	M00056217A:H11	UC2-NormColon
1391	578718	1743.K04.gz43_221534	F	M00056227D:A02	UC2-NormColon
1392	649272	1743.N04.gz43_221537	F	M00056232C:E06	UC2-NormColon
1393	648351	1743.D05.gz43_221543	F	M00056218C:C02	UC2-NormColon
1394	618670	1743.I06.gz43_221564	F	M00056225B:A11	UC2-NormColon
1395	706245	1743.P07.gz43_221587	F	M00056236A:G12	UC2-NormColon
1396	638807	1743.A08.gz43_221588	F	M00056214B:H03	UC2-NormColon
1397	562359	1743.C09.gz43_221606	_ F	M00056217B:G03	UC2-NormColon
1398	447892	1743.K09.gz43_221614	F	M00056228A:A02	UC2-NormColon
1399	502413	1743.N09.gz43_221617	F	M00056232D:C08	UC2-NormColon
1400	499517	1743.A10.gz43_221620	F	M00056214C:A04	UC2-NormColon
1401	570052	1743.G11.gz43_221642	F	M00056222A:E05	UC2-NormColon
1402	649351	1743.M11.gz43_221648	F	M00056231B:E01	UC2-NormColon
1403	650975	1743.A16.gz43_221716	F	M00056214D:F05	UC2-NormColon
1404	562625	1743.C16.gz43_221718	F	M00056217D:B07	UC2-NormColon
1405	644916	1743.E16.gz43_221720	F	M00056220A:H04	UC2-NormColon
1406	33738	1743.L16.gz43_221727	F	M00056229C:F05	UC2-NormColon
1407	645549	1743.H17.gz43_221739	F	M00056224B:F01	UC2-NormColon
1408	451487	1743.M21.gz43 221808	F	M00056232B:A11	UC2-NormColon
1409	448780	1743.C22.gz43 221814	F	M00056218A:G11	UC2-NormColon
1410	648816	1743.O22.gz43 221826	F	M00056235A:C12	UC2-NormColon
1411	648989	1743.G23.gz43 221834	F	M00056223A:B11	UC2-NormColon
1412	453470	1743.H23.gz43 221835	F	M00056224D:E08	UC2-NormColon
1413	649030	1743.M23.gz43 221840	F	M00056232B:C04	UC2-NormColon
1414	442347	1743.G24.gz43 221850	F	M00056223A:C03	UC2-NormColon
1415	645271	1744.001.gz43 221874	F	M00056256A:C02	UC2-NormColon
1416	645872	1744.J02.gz43 221885	F	M00056249A;H07	UC2-NormColon
1417	651051	1744.N02.gz43 221889	F	M00056254A:F04	UC2-NormColon
1418	650463	1744.F03.gz43_221897	F	M00056244A;A01	UC2-NormColon
1419	506719	1744.J03.gz43 221901	F	M00056249B;C02	UC2-NormColon
1420	648989	1744.E04.gz43 221912	F	M00056242D:A02	UC2-NormColon
1421	703978	1744.A06.gz43 221940	F	M00056237B:D04	UC2-NormColon
1422	650661	1744.K07.gz43 221966	F	M00056250C;D05	UC2-NormColon
1423	419255	1744.H08.gz43 221979	F	M00056246C;G07	UC2-NormColon
1424	648996	1744.M08.gz43 221984	F	M00056253A:F12	UC2-NormColon
1425	451361	1744.N09.gz43 222001	F	M00056254D:C04	UC2-NormColon
1426	15296	1744.A10.gz43 222004	F	M00056237C;C05	UC2-NormColon
1427	649396	1744.C10.gz43 222006	F	M00056240B:F08	UC2-NormColon
1428	650860	1744.J10.gz43 222013	F	M00056249C:E06	UC2-NormColon

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SEO			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
1429	643327	1744.K14,gz43 222078	F	M00056251B:A12	UC2-NormColon
1430	639119	1744.M14.gz43 222080	F	M00056253B:H04	UC2-NormColon
1431	447272	1744.C15.gz43 222086	F	M00056240D:H07	UC2-NormColon
1432	452646	1744.K15.gz43 222094	F	M00056251B:B07	UC2-NormColon
1433	637387	1744.G17.gz43 222122	F	M00056246A:E01	UC2-NormColon
1434	162981	1744.H17.gz43 222123	F	M00056247A:D02	UC2-NormColon
1435	457221	1744.H18.gz43 222139	F	M00056247A:F07	UC2-NormColon
1436	703217	1744.M18.gz43_222144	F	M00056253C:C04	UC2-NormColon
1437	559324	1744.N18.gz43 222145	F	M00056255B:F09	UC2-NormColon
1438	707483	1744.A19.gz43 222148	F	M00056238A:A03	UC2-NormColon
1439	464091	1744.P19.gz43 222163	F	M00056258A:A11	UC2-NormColon
1440	645472	1744.N20.gz43_222103	F	M00056255C:C04	UC2-NormColon
1441	650018	1744.H21.gz43 222177	F	M00056247B:C11	UC2-NormColon
1442	651073	1744.E22.gz43 222200	F	M00056247B:C11	UC2-NormColon
1443	641874	1744.M23.gz43 222224	F	M00056253D:G04	UC2-NormColon
1444	610269	1744.P23.gz43 222227	F		
1445	650463	1744.F24.gz43 222233	F	M00056258A:E10 M00056245A:D11	UC2-NormColon UC2-NormColon
1446	650919	1744.F24.gz43_222236	F		
1447	586794	1744.K24.gz43_222238	F	M00056249A;F11	UC2-NormColon
1447	642200	1753.K02.gz43 222270	F	M00056251C:H04 M00056273A:A07	UC2-NormColon
1449	645271		F		UC2-NormColon
1449	528162	1753.C03.gz43_222278 1753.J04.gz43_222301	F	M00056261A:F12	UC2-NormColon
1451	648609			M00056271C:C06	UC2-NormColon
1451	213	1753.N05.gz43_222321	F	M00056277B:G05	UC2-NormColon
1452		1753.D06.gz43_222327 1753.E06.gz43_222328	F	M00056262D:G08	UC2-NormColon
1454	648063			M00056264B:C03	UC2-NormColon
	647828	1753.I06.gz43_222332	F	M00056270A:B09	UC2-NormColon
1455	476380	1753.L06.gz43_222335	F	M00056274D:E08	UC2-NormColon
1456	649262	1753.D10.gz43_222391	F	M00056263A:D08	UC2-NormColon
1457	648590	1753.A12.gz43_222420	F	M00056258D:H06	UC2-NormColon
1458	557857	1753.I12.gz43_222428	F	M00056270B:F02	UC2-NormColon
1459	649349	1753.P13.gz43_222451	F	M00056280B:E09	UC2-NormColon
1460	460245	1753.B14.gz43_222453	F	M00056260C;E11	UC2-NormColon
1461	648442	1753.M14.gz43_222464	F	M00056276A:F05	UC2-NormColon
1462	646609	1753.E15.gz43_222472	F	M00056265A:E01	UC2-NormColon
1463	562173	1753.G15.gz43_222474	F	M00056267D:B09	UC2-NormColon
1464	555021	1753.A17.gz43_222500	F	M00056259A:E08	UC2-NormColon
1465	486452	1753.D17.gz43_222503	F	M00056263B:E10	UC2-NormColon
1466	507066	1753.M17.gz43_222512	F	M00056276D:B12	UC2-NormColon
1467	649592	1753.O18.gz43_222530	F	M00056279B:G06	UC2-NormColon
1468	268336	1753.C23.gz43_222598	F	M00056262B:B08	UC2-NormColon
1469	509202	1753.P23.gz43_222611	F	M00056280D:C06	UC2-NormColon
1470	459914	1753.A24.gz43_222612	F	M00056259C:E03	UC2-NormColon

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T	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
1471	466894	1753.I24.gz43_222620	F	M00056271A:G04	UC2-NormColon
1472	647839	1754.D07.gz43_222727	F	M00056285B:B01	UC2-NormColon
1473	644862	1754.I07.gz43_222732	F	M00056292D:C06	UC2-NormColon
1474	492587	1754.K.07.gz43_222734	F	M00056295C:F12	UC2-NormColon
1475	651057	1754.M07.gz43_222736	F	M00056298A;H02	UC2-NormColon
1476	554276	1754.E10.gz43_222776	F	M00056286C:H08	UC2-NormColon
1477	643984	1754.N10.gz43_222785	F	M00056299C:F06	UC2-NormColon
1478	646060	1754.H12.gz43_222811	F	M00056291B:G01	UC2-NormColon
1479	646387	1754.K12.gz43_222814	F	M00056296A:A07	UC2-NormColon
1480	462986	1754.N12.gz43_222817	F	M00056299D:A02	UC2-NormColon
1481	598471	1754.H19.gz43_222923	F	M00056291C:C11	UC2-NormColon
1482	449882	1754.I19.gz43_222924	F	M00056293B:F02	UC2-NormColon
1483	555998	1754.F20.gz43_222937	F	M00056289B:E05	UC2-NormColon
1484	419801	1754.J20.gz43_222941	F	M00056294B:G06	UC2-NormColon
1485	647462	1754.E21.gz43_222952	F	M00056287C:H08	UC2-NormColon
1486	648524	1754.F21.gz43_222953	F	M00056289B:F09	UC2-NormColon
1487	462865	1754.M22.gz43_222976	F	M00056298D:G01	UC2-NormColon
1488	645844	1754.O22.gz43_222978	F	M00056301C:H08	UC2-NormColon
1489	556497	1754.C23.gz43_222982	F	M00056284C:A11	UC2-NormColon
1490	467709	1754.L23.gz43_222991	F	M00056297D:B10	UC2-NormColon
1491	559857	1754.D24.gz43_222999	F	M00056286B:A04	UC2-NormColon
1492	651115	1754.N24.gz43_223009	F	M00056300B:A11	UC2-NormColon
1493	644660	1755.D05.gz43_223079	F	M00056307A:F07	UC2-NormColon
1494	647248	1755.N05.gz43_223089	F	M00056329C:B09	UC2-NormColon
1495	524261	1755.B06.gz43_223093	F	M00056305A:A02	UC2-NormColon
1496	556444	1755.K06.gz43_223102	F	M00056320C:E08	UC2-NormColon
1497	639378	1755.M08.gz43_223136	F	M00056326C:G07	UC2-NormColon
1498	460929	1755.O09.gz43_223154	F	M00056333A:E09	UC2-NormColon
1499	642260	1755.K10.gz43_223166	F	M00056320D:F07	UC2-NormColon
1500	635062	1755.I11.gz43_223180	F	M00056319A:A11	UC2-NormColon
1501	644880	1755.B14.gz43_223221	F	M00056305B:F09	UC2-NormColon
1502	452075	1755.N14.gz43_223233	F	M00056330B:C02	UC2-NormColon
1503	557052	1755.E15.gz43_223240	F	M00056310A:C10	UC2-NormColon
1504	569195	1755.K15.gz43_223246	F	M00056323A:G03	UC2-NormColon
1505	444222	1755.M16.gz43_223264	F	M00056328B:E07	UC2-NormColon
1506	467364	1755.A17.gz43_223268	F	M00056304A:H04	UC2-NormColon
1507	446910	1755.A18.gz43_223284	F	M00056304B:C05	UC2-NormColon
1508	447126	1755.M19.gz43_223312	F	M00056328D:B01	UC2-NormColon
1509	650639	1755.D20.gz43_223319	F	M00056308A:C01	UC2-NormColon
1510	641681	1755.B22.gz43_223349	F	M00056305D:E08	UC2-NormColon
1511	606129	1755.P24.gz43_223395	F	M00056338B:B08	UC2-NormColon
1512	448412	1766.A01.gz43_224099	F	M00042346A:B04	UC2-PrimColon

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Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
1513	485237	1766.E01.gz43 224103	F	M00042432D:G02	UC2-PrimColon
1514	479131	1766.G01.gz43_224105	F	M00042437B:A11	UC2-PrimColon
1515	481457	1766.H03.gz43_224138	F	M00042439A:D09	UC2-PrimColon
1516	448892	1766.A09.gz43_224227	F	M00042346C:G12	UC2-PrimColon
1517	477797	1766.E13.gz43_224295	F	M00042433C:H05	UC2-PrimColon
1518	450212	1766,I19.gz43_224395	F	M00042462A:F12	UC2-PrimColon
1519	3781	1766.J21.gz43_224428	F	M00042464A:F06	UC2-PrimColon
1520	447377	1766,M22.gz43 224447	F	M00042470B:H11	UC2-PrimColon
1521	453606	1767.H01.gz43_224490	F	M00042764B:B10	UC2-PrimColon
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1553	448260	1767.O15.gz43_224721	F	M00042778B:B07	UC2-PrimColon
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Table 2

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1559 485907 1767.C17.gz43_224741 F M00042755B:H11 UC2-PrimColon 1560 444830 1767.F17.gz43_224744 F M00042776D:D10 UC2-PrimColon 1561 448152 1767.F17.gz43_224754 F M00042776D:D10 UC2-PrimColon 1562 474823 1767.F17.gz43_224754 F M00042776D:D10 UC2-PrimColon 1562 474823 1767.F17.gz43_224754 F M00042776D:D10 UC2-PrimColon 1563 433024 1767.E18.gz43_224758 F M00042767B:B10 UC2-PrimColon 1564 479880 1767.H18.gz43_224768 F M00042767B:B10 UC2-PrimColon 1565 477399 1767.N18.gz43_224768 F M00042751D:G09 UC2-PrimColon 1566 485441 1767.A19.gz43_224771 F M00042751D:G09 UC2-PrimColon 1566 485441 1767.A19.gz43_224771 F M00042751D:G09 UC2-PrimColon 1567 485183 1767.B19.gz43_224772 F M00042755C:A10 UC2-PrimColon 1568 452104 1767.C19.gz43_224773 F M00042755C:A10 UC2-PrimColon 1570 447218 1767.L20.gz43_224980 F M00042775C:C12 UC2-PrimColon 1571 472119 1767.N20.gz43_224800 F M00042775C:C12 UC2-PrimColon 1572 482868 1767.D12.gz43_224811 F M00042776B:B10 UC2-PrimColon 1574 484964 1767.L22.gz43_224827 F M00042767B:B10 UC2-PrimColon 1574 484964 1767.L22.gz43_224827 F M00042758B:G04 UC2-PrimColon 1575 448325 1767.D23.gz43_224839 F M0004276B:G04 UC2-PrimColon 1576 483987 1767.D23.gz43_224839 F M0004276B:G04 UC2-PrimColon 1577 475797 1767.D23.gz43_224849 F M0004276B:G04 UC2-PrimColon 1578 448233 1778.D01.gz43_225256 F M00042834A:B11 UC2-PrimColon 1584 449403 1778.N02.gz43_225309 F M00042850D:P05 UC2-PrimColon 1584 449403 1778.N02.gz43_225309 F M00042850D:P05 UC2-PrimColon 1584 449604 1778.N03.gz43_225309 F M00042850D:P05 UC2-PrimCol	1557	449592	1767.L16.gz43_224734	F	M00042771D:F02	UC2-PrimColon
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1563	1561	448152	1767.N17.gz43_224752	F	M00042776D:D01	UC2-PrimColon
1564 479880 1767.118.gz43_224763 F M00042767A:B10 UC2-PrimColon 1565 4875441 1767.N18.gz43_224767 F M00042775D:G10 UC2-PrimColon 1566 485441 1767.N19.gz43_224771 F M00042751D:G09 UC2-PrimColon 1567 485183 1767.B19.gz43_224772 F M00042751D:G10 UC2-PrimColon 1568 452104 1767.C19.gz43_224773 F M00042753C:G12 UC2-PrimColon 1569 449275 1767.A20.gz43_224787 F M00042755C:A10 UC2-PrimColon 1570 447218 1767.L20.gz43_224787 F M00042753C:B04 UC2-PrimColon 1571 472119 1767.N20.gz43_224800 F M000427762A:B04 UC2-PrimColon 1572 482868 1767.121.gz43_224811 F M00042756A:G02 UC2-PrimColon 1573 4484964 1767.122.gz43_224824 F M00042767B:G09 UC2-PrimColon 1574 4484964 1767.122.gz43_224823 F M0004276B:G04 UC2-PrimColon 1575 448325 1767.D23.gz43_224833 F M0004276B:G04 UC2-PrimColon 1576 4484987 1767.023.gz43_224843 F M0004276B:G09 UC2-PrimColon 1577 475797 1767.023.gz43_224849 F M0004276B:G09 UC2-PrimColon 1578 477757 1767.023.gz43_224866 F M0004276B:G09 UC2-PrimColon 1579 448233 1778.D01.gz43_225308 F M000428842A:B12 UC2-PrimColon 1580 475111 1778.B02.gz43_225288 F M00042881A:B02 UC2-PrimColon 1581 449403 1778.N02.gz43_225288 F M00042881A:B02 UC2-PrimColon 1582 449604 1778.O02.gz43_225308 F M00042881A:B02 UC2-PrimColon 1583 447983 1778.D02.gz43_225308 F M00042850D:F02 UC2-PrimColon 1584 484091 1778.O03.gz43_225308 F M00042851B:D06 UC2-PrimColon 1585 485841 1778.L05.gz43_225309 F M00042851B:D06 UC2-PrimColon 1589 447983 1778.B03.gz43_225309 F M00042850D:F05 UC2-PrimColon 1580 47518 1778.L05.gz43_225309 F M00042855D:A12 UC2-PrimColon 1580 47518 1778.L05.gz43_225309 F M00042855D:A12 UC2-PrimColon 1590 725641 1778.L05.gz43_225309 F M00042855D:A12 UC2-PrimColon 1590 475641 1778.L05.gz43_225309 F M00042855D:A12 UC2-PrimC	1562	474823	1767.P17.gz43_224754	F	M00042779D:E06	UC2-PrimColon
1565 477399 1767.N18.gz43_224768 F M00042776D:G10 UC2-PrimColon 1566 485441 1767.A19.gz43_224771 F M000427351D:G09 UC2-PrimColon 1567 485183 1767.B19.gz43_224772 F M000427351D:G12 UC2-PrimColon 1568 452104 1767.C19.gz43_224773 F M00042735C:G12 UC2-PrimColon 1569 449275 1767.A20.gz43_224787 F M00042735C:A10 UC2-PrimColon 1570 447218 1767.L20.gz43_224787 F M00042735C:C12 UC2-PrimColon 1570 447218 1767.L20.gz43_224980 F M00042775C:C12 UC2-PrimColon 1571 472119 1767.N20.gz43_224800 F M000427767B:B10 UC2-PrimColon 1572 482868 1767.I21.gz43_224811 F M0004276763C02 UC2-PrimColon 1573 455143 1767.P22.gz43_224824 F M0004276763C02 UC2-PrimColon 1574 484964 1767.I22.gz43_224824 F M0004276763C02 UC2-PrimColon 1575 448325 1767.D23.gz43_224838 F M00042758B:G00 UC2-PrimColon 1576 484987 1767.D23.gz43_224838 F M00042758B:G00 UC2-PrimColon 1576 484987 1767.D23.gz43_224843 F M00042758B:G00 UC2-PrimColon 1577 475797 1767.P24.gz43_224866 F M0004278D:F11 UC2-PrimColon 1578 447931 1778.D01.gz43_225254 F M00042845A12 UC2-PrimColon 1584 449403 1778.N02.gz43_225258 F M00042845A12 UC2-PrimColon 1582 449604 1778.002.gz43_225288 F M00042859B:D06 UC2-PrimColon 1582 449604 1778.002.gz43_225288 F M00042850D:P05 UC2-PrimColon 1584 484091 1778.D03.gz43_225288 F M00042850D:P05 UC2-PrimColon 1585 489000 1778.N03.gz43_225280 F M00042850D:P05 UC2-PrimColon 1584 484091 1778.L03.gz43_225309 F M00042850D:P05 UC2-PrimColon 1585 485841 1778.L03.gz43_225309 F M00042850D:P05 UC2-PrimColon 1586 475181 1778.L03.gz43_225309 F M00042850D:P05 UC2-PrimColon 1589 447983 1778.L03.gz43_225309 F M00042850D:P05 UC2-PrimColon 1589 447983 1778.L03.gz43_225309 F M00042850D:P05 UC2-PrimColon 1590 484789 1778.L03.gz43_225309 F M00042850D:D05 UC2-PrimCo	1563	453024	1767.E18.gz43 224759	F.	M00042760B:E12	UC2-PrimColon
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1567	1565	477399	1767.N18.gz43 224768	F	M00042776D:G10	UC2-PrimColon
1568 452104 1767.C19.gz43_224773 F M00042755C:A10 UC2-PrimColon 1569 449275 1767.A20.gz43_224787 F M00042752A:B04 UC2-PrimColon 1570 447218 1767.L20.gz43_224898 F M00042775C:C12 UC2-PrimColon 1571 472119 1767.N20.gz43_224800 F M000427767B:B10 UC2-PrimColon 1572 482868 1767.I21.gz43_224811 F M0004276763C02 UC2-PrimColon 1573 455134 1767.P22.gz43_224824 F M0004276763C02 UC2-PrimColon 1574 484964 1767.I22.gz43_224824 F M0004276763C02 UC2-PrimColon 1575 448325 1767.D23.gz43_224827 F M000427578:G04 UC2-PrimColon 1576 448947 1767.123.gz43_224834 F M000427578:G09 UC2-PrimColon 1577 475797 1767.D23.gz43_224834 F M000427578:G09 UC2-PrimColon 1578 448494 1767.123.gz43_224843 F M0004278D:F11 UC2-PrimColon 1579 44823 1778.D01.gz43_225254 F M0004278D:H10 UC2-PrimColon 1580 475111 1778.B02.gz43_225254 F M00042848-B12 UC2-PrimColon 1581 449403 1778.N02.gz43_225280 F M0004288A:E02 UC2-PrimColon 1582 449604 1778.002.gz43_225288 F M0004288D:P60 UC2-PrimColon 1583 447983 1778.F03.gz43_225309 F M00042850D:P60 UC2-PrimColon 1584 484091 1778.003.gz43_225309 F M0004285D:P60 UC2-PrimColon 1585 485090 1778.N04.gz43_225309 F M00042851B:H0 UC2-PrimColon 1586 4551811 1778.K04.gz43_225330 F M00042851B:H0 UC2-PrimColon 1587 4487983 1778.E07.gz43_225351 F M00042851B:H0 UC2-PrimColon 1588 4476256 1778.C07.gz43_225351 F M00042851B:H0 UC2-PrimColon 1589 447983 1778.E07.gz43_225351 F M00042851A:D0 UC2-PrimColon 1590 484789 1778.L09.gz43_225351 F M00042851A:D0 UC2-PrimColon 1591 484789 1778.L09.gz43_225350 F M00042851A:D0 UC2-PrimColon 1592 449110 1778.L09.gz43_225350 F M00042851A:D0 UC2-PrimColon 1592 449110 1778.L09.gz43_225350 F M00042850:D-12 UC2-PrimColon 1593 451368 1778.C09.gz	1566	485441	1767,A19.gz43 224771	F	M00042751D:G09	UC2-PrimColon
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1571 472119 1767.N20.gz43 224800 F M00042777A:B10 UC2-PrimColon 1572 482868 1767.I21.gz43 224811 F M00042767B:B10 UC2-PrimColon 1573 455143 1767.P22.gz43 224824 F M00042767B:G02 UC2-PrimColon 1574 484964 1767.I22.gz43 224827 F M00042767B:G04 UC2-PrimColon 1575 448325 1767.D23.gz43 224827 F M00042757B:G04 UC2-PrimColon 1575 448947 1767.I23.gz43 224824 F M00042757B:G06 UC2-PrimColon 1576 448947 1767.D23.gz43 224849 F M00042757B:G09 UC2-PrimColon 1577 475797 1767.P24.gz43 224866 F M00042780A:H05 UC2-PrimColon 1578 447757 1767.P24.gz43 224866 F M00042780A:H05 UC2-PrimColon 1579 44823 1778.D01.gz43 225254 F M00042840A:B12 UC2-PrimColon 1580 475111 1778.B02.gz43 225268 F M00042859B:D06 UC2-PrimColon 1581 449403 1778.N02.gz43 225288 F M00042859B:D06 UC2-PrimColon 1582 449604 1778.002.gz43 225288 F M00042850D:P02 UC2-PrimColon 1583 4447983 1778.F03.gz43 225288 F M00042860D:P02 UC2-PrimColon 1584 484091 1778.003.gz43 225309 F M00042850D:P05 UC2-PrimColon 1585 480900 1778.N03.gz43 225309 F M00042850D:P05 UC2-PrimColon 1586 551811 1778.K04.gz43 225309 F M00042851B:H06 UC2-PrimColon 1586 476256 1778.C07.gz43 225351 F M00042851B:H06 UC2-PrimColon 1589 447983 1778.E07.gz43 225351 F M00042851A:D0 UC2-PrimColon 1589 447983 1778.L07.gz43 225351 F M00042855D:A12 UC2-PrimColon 1590 484789 1778.L09.gz43 225309 F M00042855D:A12 UC2-PrimColon 1590 484789 1778.L09.gz43 225351 F M00042855D:A12 UC2-PrimColon 1590 484789 1778.L09.gz43 225390 F M00042855D:A12 UC2-PrimColon 1590 484789 1778.L09.gz43 225390 F M00042850D:A10 UC2-PrimColon 1590 484789 1778.L09.gz43 225390 F M00042850D:A10 UC2-PrimColon 1590 481368 1778.L09.gz43 225390 F M00042850D:A10 UC2-PrimColon 1590 481368 1	1569	449275	1767,A20,gz43 224787	F	M00042752A:B04	UC2-PrimColon
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1578				F		
1579	1578			F		
1580 475111 1778.B02.gz43_225268 F M00042838A:E02 UC2-PrimColon	1579	448233		F	M00042842A:B12	
1581 449403 1778.N02.gz43 225280 F M00042859B:D06 UC2-PrimColon 1582 449604 1778.002.gz43 225281 F M00042850D:P02 UC2-PrimColon 1583 447983 1778.F03.gz43 225288 F M000428450D:P02 UC2-PrimColon 1584 484091 1778.003.gz43 225297 F M00042850D:P05 UC2-PrimColon 1585 485090 1778.104.gz43 225309 F M00042853B:P06 UC2-PrimColon 1586 551811 1778.K04.gz43 225309 F M00042854B:B11 UC2-PrimColon 1587 485841 1778.L05.gz43 225309 F M00042854B:B16 UC2-PrimColon 1588 447625 1778.C07.gz43 225349 F M00042854B:B16 UC2-PrimColon 1589 447983 1778.E07.gz43 225351 F M00042844D:D02 UC2-PrimColon 1590 725641 1778.K07.gz43 225351 F M00042845A:D05 UC2-PrimColon 1591 484789 1778.L08.gz43 225374 F M00042855C:G11 UC2-PrimColon 1592 449110 1778.L09.gz43 225390 F M00042851A:P04 UC2-PrimColon 1593 4419110 1778.L09.gz43 225390 F M00042861A:P04 UC2-PrimColon 1594 449521 1778.P09.gz43 225394 F M00042861A:P04 UC2-PrimColon 1595 727875 1778.G10.gz43 225304 F M00042861A:P04 UC2-PrimColon 1595 727875 1778.G10.gz43 225304 F M00042861A:P04 UC2-PrimColon	1580			F		
1582 449604 1778,002,gz43 225281 F M00042860D:F02 UC2-PrimColon	1581	449403		F		
1583 447983 1778,F03,gz43,225288 F M00042845C;A09 UC2-PrimColon 1584 484091 1778,O03,gz43,225297 F M00042860D;F05 UC2-PrimColon 1585 489090 1778,D04,gz43,225308 F M0004285CD;C06 UC2-PrimColon 1586 551811 1778,K04,gz43,225309 F M00042854A;B11 UC2-PrimColon 1587 485841 1778,L05,gz43,225326 F M00042855B;H06 UC2-PrimColon 1588 476256 1778,C07,gz43,225339 F M00042835D;H02 UC2-PrimColon 1589 447983 1778,E07,gz43,225351 F M00042834A;D02 UC2-PrimColon 1590 725641 1778,K07,gz43,225374 F M00042855C;G11 UC2-PrimColon 1591 484789 1778,L09,gz43,225374 F M00042855D;A12 UC2-PrimColon 1592 449110 1778,L09,gz43,225390 F M00042855D;A12 UC2-PrimColon 1593 451368 1778,D99,gz43,225393 F M00042854A;D04 UC2-PrimColon 1594 449521 1778,P09,gz43,225394 F M00042861A;P04 UC2-PrimColon 1595 727875 1778,F01,gz43,225401 F M00042847A;P04 UC2-PrimColon 1595 727875 1778,F01,gz43,225394 F M00042861C;E02 UC2-PrimColon 1595 727875 1778,F01,gz43,225394 F M00042847A;P04 UC2-PrimColon				F	M00042860D:F02	
1584 484091 1778.003 gz/3 225207 F M00042850D:P65 UC2-PrimColon 1585 480900 1778.104 gz/3 225308 F M00042852B:C06 UC2-PrimColon 1586 551811 1778.K04 gz/3 225309 F M00042854A:B11 UC2-PrimColon 1587 485841 1778.L05 gz/3 225326 F M00042855B:H06 UC2-PrimColon 1589 447983 1778.E07 gz/3 225351 F M00042834A:D02 UC2-PrimColon 1590 725641 1778.L07 gz/3 225374 F M00042834A:D05 UC2-PrimColon 1591 484789 1778.L08 gz/3 225390 F M00042855C:G11 UC2-PrimColon 1592 449110 1778.L09 gz/3 225390 F M00042851A:P04 UC2-PrimColon 1593 451368 1778.P09 gz/3 225394 F M00042861A:P04 UC2-PrimColon 1594 449521 1778.P09 gz/3 225394 F M00042861A:P04 UC2-PrimColon 1595 727875 1778.G10 gz/3 225401 F M00042847A:P04 UC2-PrimColon	1583			F	M00042845C:A09	
1585 480900 1778.104.gr43 225308 F M00042852B:C06 UC2-PrimColon 1586 551811 1778.K04.gr43 225309 F M00042854A:B11 UC2-PrimColon 1587 485841 1778.K05.gr43 225326 F M00042854B:B16 UC2-PrimColon 1588 476256 1778.C07.gr43 225349 F M00042855B:H06 UC2-PrimColon 1589 447983 1778.E07.gr43 225351 F M00042854A:D02 UC2-PrimColon 1590 725641 1778.K07.gr43 225357 F M00042854A:D05 UC2-PrimColon 1591 4347789 1778.K07.gr43 225374 F M00042855D:A12 UC2-PrimColon 1592 449110 1778.L09.gr43 225390 F M00042855D:A12 UC2-PrimColon 1593 451368 1778.09.gr43 225394 F M00042861A:P04 UC2-PrimColon 1594 449521 1778.P09.gr43 225394 F M00042862C:B02 UC2-PrimColon 1595 727875 1778.G10.gr43 225401 F M00042862C:B02 UC2-PrimColon 1595 727875 1778.G10.gr43 225401 F M00042874:P04 UC2-PrimColon	1584	484091		F		
1586 551811 1778.K04.gz43 225309 F M00042854A:B11 UC2-PrimColon 1587 485841 1778.L05.gz43 225326 F M00042855B:H06 UC2-PrimColon 1588 476256 1778.C07.gz43 225349 F M000428391-PO2 UC2-PrimColon 1589 447983 1778.E07.gz43 225351 F M000428391-PO2 UC2-PrimColon 1590 725641 1778.K07.gz43 225357 F M00042834A:D05 UC2-PrimColon 1591 484789 1778.L09.gz43 225374 F M00042854C:D1 UC2-PrimColon 1592 449110 1778.L09.gz43 225390 F M00042855D:A12 UC2-PrimColon 1593 451368 1778.099.gz43 225393 F M000428561A:P04 UC2-PrimColon 1594 449521 1778.P09.gz43 225394 F M000428502:E02 UC2-PrimColon 1595 727875 1778.G10.gz43 225301 F M00042847A:P04 UC2-PrimColon	1585			F	M00042852B:C06	
1587 485841 1778.L05.gz43.225326 F M00042855B:H06 UC2-PrimColon 1588 476256 1778.C07.gz43.225349 F M00042845102 UC2-PrimColon 1589 447983 1778.E07.gz43.225351 F M000428451020 UC2-PrimColon 1590 725641 1778.L08.gz43.225374 F M000428451005 UC2-PrimColon 1591 448719 1778.L08.gz43.225370 F M00042855C:G1 UC2-PrimColon 1592 449110 1778.L09.gz43.225390 F M000428510-L12 UC2-PrimColon 1593 451368 1778.D09.gz43.225393 F M00042861A:F04 UC2-PrimColon 1594 449521 1778.P09.gz43.225394 F M00042860C:E02 UC2-PrimColon 1595 727875 1778.G10.gz43.225401 F M00042847A:F04 UC2-PrimColon	1586	551811		F		
1588 476256 1778.C07.gz43_225349 F M00042839D:F02 UC2-PrimColon 1589 447983 1778.E07.gz43_225351 F M00042834A:D02 UC2-PrimColon 1590 725641 1778.K07.gz43_225357 F M00042854A:D05 UC2-PrimColon 1591 484789 1778.L08.gz43_225374 F M00042855C:G11 UC2-PrimColon 1592 449110 1778.L09.gz43_225390 F M00042855D:A12 UC2-PrimColon 1593 451368 1778.009.gz43_225393 F M00042861A:F04 UC2-PrimColon 1594 449521 1778.P09.gz43_225394 F M00042862C:E02 UC2-PrimColon 1595 727875 1778.G10.gz43_225401 F M00042847A:F04 UC2-PrimColon						
1589 447983 1778_E07_gz43_225351 F M00042844A:D02 UC2-PrimColon 1590 725641 1778_K07_gz43_225357 F M00042854A:D05 UC2-PrimColon 1591 484789 1778_K08_gz43_225374 F M00042855C;G1 UC2-PrimColon 1592 449110 1778_L09_gz43_225390 F M00042855D:A12 UC2-PrimColon 1593 451368 1778_099_gz43_225393 F M00042861A:P04 UC2-PrimColon 1594 449521 1778_P09_gz43_225394 F M00042862C:E02 UC2-PrimColon 1595 727875 1778_G10_gz43_225401 F M00042847A:P04 UC2-PrimColon						
1590 725641 1778.K07.gz43 225357 F M00042854A:D05 UC2-PrimColon 1591 484789 1778.L08.gz43 225374 F M00042855C:G11 UC2-PrimColon 1592 449110 1778.L09.gz43 225390 F M00042855D:A12 UC2-PrimColon 1593 451368 1778.009.gz43 225393 F M00042861A:F04 UC2-PrimColon 1594 449521 1778.P09.gz43 225394 F M0004286C:E02 UC2-PrimColon 1595 727875 1778.G10.gz43 225401 F M00042847A:F04 UC2-PrimColon 1595 1778.G10.gz43 1778.G						
1591 484789 1778.L08.gz43_225374 F M00042855C:G11 UC2-PrimColon 1592 449110 1778.L09.gz43_225390 F M00042855D:A12 UC2-PrimColon 1593 451368 1778.009.gz43_225393 F M00042861A:F04 UC2-PrimColon 1594 449521 1778.P09.gz43_225394 F M00042862C:B02 UC2-PrimColon 1595 727875 1778.G10.gz43_225401 F M00042847A:F04 UC2-PrimColon						
1592 449110 1778.L09_gz43_225390 F M00042855D:A12 UC2-PrimColon 1593 451368 1778.099_gz43_225393 F M00042861A:P04 UC2-PrimColon 1594 449521 1778.P09_gz43_225394 F M00042862C:E02 UC2-PrimColon 1595 727875 1778.G10_gz43_225401 F M00042847A:P04 UC2-PrimColon						
1593 451368 1778.009.gx43_225393 F M00042861A:F04 UC2-PrimColon 1594 449521 1778.P09.gx43_225394 F M00042862C:E02 UC2-PrimColon 1595 727875 1778.G10.gx43_225401 F M00042847A:F04 UC2-PrimColon						
1594 449521 1778.P09.gz43 225394 F M00042862C:E02 UC2-PrimColon 1595 727875 1778.G10.gz43 225401 F M00042847A:F04 UC2-PrimColon						
1595 727875 1778.G10.gz43_225401 F M00042847A:F04 UC2-PrimColon				_		
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Table 2

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1598	481664	1778.N11.gz43 225424	F	M00042859D:D12	UC2-PrimColon
1599	448230	1778.E12.gz43 225431	F	M00042844C;B03	UC2-PrimColon
1600	485029	1778.H12.gz43_225434	F	M00042849B:G06	UC2-PrimColon
1601	483274	1778.I12.gz43_225435	F	M00042851A:E11	UC2-PrimColon
1602	644190	1778.J12.gz43_225436	F	M00042853A:G03	UC2-PrimColon
1603	725266	1778.K12.gz43_225437	F	M00042854B:F05	UC2-PrimColon
1604	480005	1778.L12.gz43_225438	F	M00042855D:B12	UC2-PrimColon
1605	484841	1778.M12.gz43_225439	F	M00042857C:G05	UC2-PrimColon
1606	480641	1778,N12.gz43_225440	F	M00042860A:C01	UC2-PrimColon
1607	481319	1778.O13.gz43_225457	F	M00042861C:C03	UC2-PrimColon
1608	387077	1778.C14.gz43_225461	F	M00042840C:B12	UC2-PrimColon
1609	456737	1778.L14.gz43_225470	F	M00042855D:D06	UC2-PrimColon
1610	479084	1778,M14.gz43_225471	F	M00042858A:A08	UC2-PrimColon
1611	484468	1778.O15.gz43_225489	F	M00042861C:F07	UC2-PrimColon
1612	450673	1778.C18.gz43_225525	F	M00042841B:H05	UC2-PrimColon
1613	474597	1778.D18.gz43_225526	F	M00042843A:E07	UC2-PrimColon
1614	477593	1778.E18.gz43_225527	F	M00042844D:H02	UC2-PrimColon
1615	452990	1778.I18.gz43_225531	F	M00042851D:B08	UC2-PrimColon
1616	449407	1778.O18.gz43_225537	F	M00042861D:D09	UC2-PrimColon
1617	448104	1778.E19.gz43_225543	F	M00042845A:B05	UC2-PrimColon
1618	453856	1778.D22.gz43_225590	F	M00042843B:F05	UC2-PrimColon
1619	475942	1778.B23.gz43_225604	F	M00042839B:F05	UC2-PrimColon
1620	639991	1778.F23.gz43_225608	F	M00042846C:D09	UC2-PrimColon
1621	480640	1778.023.gz43_225617	F	M00042862A;C01	UC2-PrimColon
1622	472307	1778.C24.gz43_225621	F	M00042842A:B04	UC2-PrimColon
1623	733806	1780,I01.gz43_226027	F	M00056361A;C01	UC2-PrimColon
1624	732150	1780.E02.gz43_226039	F	M00056356C:H07	UC2-PrimColon
1625	724907	1780.K02.gz43_226045	F	M00056363A:C06	UC2-PrimColon
1626	724792	1780.L02.gz43 226046	F	M00056364A:B04	UC2-PrimColon
1627	736210	1780.B04.gz/43_226068	F	M00056353C:E12	UC2-PrimColon
1628	623978	1780.G05.gz43_226089	F	M00056359A:C08	UC2-PrimColon
1629	600856	1780.I05.gz43_226091	F	M00056361A:H06	UC2-PrimColon
1630	364334	1780.K05.gz43_226093	F	M00056363A:E10	UC2-PrimColon
1631	661194	1780.P05.gz43_226098	F	M00056368B;H09	UC2-PrimColon
1632	416326	1780.G06.gz43_226105	F	M00056359A:F06	UC2-PrimColon
1633	727129	1780.D07.gz43_226118	F	M00056355D:A05	UC2-PrimColon
1634	560859	1780.O07.gz43_226129	F	M00056367C:D06	UC2-PrimColon
1635	482831	1780.P07.gz43_226130	F	M00056368C:E07	UC2-PrimColon
1636	503862	1780.L08.gz43_226142	F	M00056364C:B01	UC2-PrimColon
1637	468467	1780.D09.gz43 226150	F	M00056355D:B08	UC2-PrimColon
1638	594994	1780.A10.gz43_226163	F	M00056352D:H02	UC2-PrimColon

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Table 2

1663 186372 1789.MO2.gz43 226431 F M00056384D:A04 UC2-PrimColor 1664 736001 1789.A03.gz43 226435 F M00056369B:B03 UC2-PrimColor 1665 487079 1789.F03.gz43 226440 F M00056374C:G03 UC2-PrimColor 1666 460284 1789.H03.gz43 226442 F M00056377A:C01 UC2-PrimColor 1667 725274 1789.I03.gz43 226443 F M00056377D:F08 UC2-PrimColor 1668 553035 1789.N03.gz43 226443 F M0005637B:B03 UC2-PrimColor 1669 724009 1789.P03.gz43 226450 F M0005638B:B03 UC2-PrimColor 1670 568484 1789.B04.gz43 226452 F M0005638B:B03 UC2-PrimColor 1671 725321 1789.L04.gz43 226452 F M0005638B:F08 UC2-PrimColor 1672 731542 1789.K05.gz43 226477 F M0005638B:F08 UC2-PrimColor 1672 731542 1789.L06.gz43 226494 F M0005638B:G08 UC2-PrimColor 1674 452806 1789.N06.gz43 226496 F M0005638B:G07 UC2-PrimColor 1675 549790 1789.L07.gz43 226510 F M0005638C:G07 UC2-PrimColor 1675 549790 1789.L07.gz43 226510 F M0005638G:A07 UC2-PrimColor 1676 730660 1789.N07.gz43 226510 F M0005638G:A11 UC2-PrimColor 1676 1780.G007.gz43 226510 F M0005638G:A11 UC2-PrimColor 1676 1780.G007		able 2			·····	
IDNO CLUSTER SEQ NAME	SEO			ODIEN		
1639		CLUSTER	SEO NAME		CI ONE ID	TIRDADV
1640						
1641 731925 1780,B11,gz43 226180 F M00056353D:D10 UC2-PrimColor 1642 480924 1780,M11,gz43 226191 F M0005636B:G10 UC2-PrimColor 1643 171298 1780,P11,gz43 226194 F M0005636B:D03 UC2-PrimColor 1644 734150 1780,F12,gz43 226200 F M0005636B:D03 UC2-PrimColor 1644 734150 1780,F12,gz43 226200 F M0005636B:D03 UC2-PrimColor 1645 528981 1780,H12,gz43 226200 F M0005636B:D09 UC2-PrimColor 1646 135364 1780,H12,gz43 226203 F M00056360B:D09 UC2-PrimColor 1647 511606 1780,J12,gz43 226203 F M0005636C:C02 UC2-PrimColor 1648 552891 1780,K14,gz43 226203 F M0005636C:C02 UC2-PrimColor 1649 730301 1780,H15,gz43 226250 F M0005636B:D03 UC2-PrimColor 1650 480005 1780,H15,gz43 226250 F M0005636B:D03 UC2-PrimColor 1651 588959 1780,L15,gz43 226250 F M0005636C:B01 UC2-PrimColor 1652 557029 1780,C16,gz43 226261 F M0005636B:D02 UC2-PrimColor 1653 553244 1780,L17,gz43 226286 F M0005636B:D02 UC2-PrimColor 1654 456535 1780,C02,gz43 226365 F M0005636B:D03 UC2-PrimColor 1655 7274039 1780,E23,gz43 226356 F M0005636B:D03 UC2-PrimColor 1656 724039 1780,E23,gz43 226375 F M0005636B:D10 UC2-PrimColor 1657 732144 1780,E23,gz43 226415 F M0005638A:CH04 UC2-PrimColor 1658 724257 1789,M01,gz43 226415 F M0005638B:D10 UC2-PrimColor 1658 724257 1789,M01,gz43 226415 F M0005638B:D10 UC2-PrimColor 1660 376342 1789,M01,gz43 226415 F M0005638B:D3 UC2-PrimColor 1661 475682 1789,M01,gz43 226415 F M0005638B:D3 UC2-PrimColor 1664 73600 1789,M03,gz43 226415 F M0005638B:D3 UC2-PrimColor 1665 487079 1789,M03,gz43 226440 F M0005638B:D3 UC2-PrimColor 1666 460284 1789,M03,gz43 226445 F M0005638B:D3 UC2-PrimColor 1666 460284 1789,M03,gz43 226445 F M0005638B:D3 UC2-PrimColor 1667 725274 1789,M03,gz43 226445 F						
1642						
1644						
1644 734150 1780.F12.g243 226200 F M00056358A:G05 UC2.PrimColor 1645 528981 1780.H12.g243 226203 F M00056360B:D09 UC2.PrimColor 1646 135364 1780.H12.g243 226203 F M00056361C:C05 UC2.PrimColor 1647 511606 1780.H12.g243 226203 F M00056362C:C02 UC2.PrimColor 1648 552891 1780.K14.g243 226237 F M00056363C:C12 UC2.PrimColor 1649 730301 1780.H14.g243 226242 F M00056363B:G05 UC2.PrimColor 1659 480005 1780.H15.g243 226250 F M00056368D:G05 UC2.PrimColor 1651 588959 1780.L15.g243 226254 F M00056366C:B01 UC2.PrimColor 1651 588959 1780.L15.g243 226254 F M00056366C:B01 UC2.PrimColor 1652 557029 1780.C16.g243 226261 F M00056358D:D02 UC2.PrimColor 1652 557029 1780.C16.g243 226261 F M00056366C:B01 UC2.PrimColor 1654 456535 1780.C02.g243 226365 F M00056363B:D02 UC2.PrimColor 1655 724774 1780.E22.g243 226365 F M00056358D:D02 UC2.PrimColor 1655 724039 1780.E22.g243 226365 F M00056358D:H03 UC2.PrimColor 1657 732144 1780.H23.g243 226415 F M00056357C:C06 UC2.PrimColor 1658 724257 1789.G01.g243 226415 F M00056357D:D12 UC2.PrimColor 1659 474346 1789.H01.g243 226415 F M0005638A:CH04 UC2.PrimColor 1660 376342 1789.H01.g243 226415 F M0005638B:A05 UC2.PrimColor 1661 475682 1789.H01.g243 226415 F M0005638B:A05 UC2.PrimColor 1662 734646 1789.H02.g243 226415 F M0005637B:C10 UC2.PrimColor 1664 756001 1789.H03.g243 226415 F M0005637B:C10 UC2.PrimColor 1664 756001 1789.H03.g243 226415 F M0005637B:C10 UC2.PrimColor 1665 487079 1789.H03.g243 226415 F M0005637B:C10 UC2.PrimColor 1666 460284 1789.H03.g243 226415 F M0005637B:C10 UC2.PrimColor 1666 460284 1789.H03.g243 226415 F M0005637B:C00 UC2.PrimColor 1667 725274 1789.H03.g243 226415 F M0005638B:B03 UC2.PrimColor 1667 725274 1789.H03.						
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1658 724257 1789.GO1.g243 226409 F M00056375D:D12 UC2-PrimColor 1659 474346 1789.MO1.g243 226415 F M00056384C:H04 UC2-PrimColor 1660 376342 1789.MO1.g243 226416 F M000563846.P08 UC2-PrimColor 1661 475682 1789.PO1.g243 226418 F M0005638A6.P08 UC2-PrimColor 1662 186372 1789.MO2.g243 226420 F M00056370B:C10 UC2-PrimColor 1663 186372 1789.MO2.g243 226431 F M00056370B:C10 UC2-PrimColor 1664 736001 1789.AO3.g243 226431 F M000563740.PO4 UC2-PrimColor 1665 487079 1789.FO3.g243 226440 F M00056374C:G08 UC2-PrimColor 1665 487079 1789.FO3.g243 226440 F M00056377A:C01 UC2-PrimColor 1666 460284 1789.H03.g243 226442 F M00056377A:C01 UC2-PrimColor 1668 553035 1789.NO3.g243 226443 F M00056376B:E03 UC2-PrimColor 1669 724009 1789.PO3.g243 226443 F M00056386A:G01 UC2-PrimColor 1669 724009 1789.PO3.g243 226452 F M00056380B:B03 UC2-PrimColor 1670 558484 1789.B04.g243 226452 F M0005638B:B03 UC2-PrimColor 1671 725321 1789.L04.g243 226462 F M0005638B:B03 UC2-PrimColor 1672 731542 1789.L04.g243 226462 F M0005638B:G03 UC2-PrimColor 1673 725448 1789.L06.g243 226494 F M0005638B:G03 UC2-PrimColor 1673 725448 1789.L06.g243 226494 F M0005638B:G07 UC2-PrimColor 1675 549790 1789.NO7.g243 226512 F M0005638B:A07 UC2-PrimColor 1675 549790 1789.NO7.g243 226512 F M0005638B:A07 UC2-PrimColor 1676 730660 1789.NO7.g243 226512 F M0005638B:A01 UC2-PrimColor 1676 730660 1789.NO7.g243 226512 F M0005638B:A11 UC2-PrimColor 1676 730660 1789.NO7.g243						
1659						
1660 376342 1789.N01.gz43 226416 F M00056386A:F08 UC2-PrimColor				_		
1661	-					
1662						
1663	_					UC2-PrimColon
1664 736001 1789.A03.gz43 226435 F M00056369B:E03 UC2-PrimColor 1665 487079 1789.P03.gz43 226440 F M00056374C:G08 UC2-PrimColor 1666 460284 1789.H03.gz43 226442 F M00056377A:C01 UC2-PrimColor 1667 725274 1789.I03.gz43 226443 F M00056377D:F08 UC2-PrimColor 1668 553035 1789.N03.gz43 226448 F M00056386A:G01 UC2-PrimColor 1669 724009 1789.P03.gz43 226450 F M00056388B:B03 UC2-PrimColor 1670 568484 1789.B04.gz43 226452 F M00056388B:B03 UC2-PrimColor 1671 725321 1789.L04.gz43 226462 F M00056383B:F08 UC2-PrimColor 1672 731542 1789.L04.gz43 226462 F M00056383B:F08 UC2-PrimColor 1673 725448 1789.L06.gz43 226494 F M0005638B:G03 UC2-PrimColor 1674 452806 1789.N06.gz43 226496 F M0005638B:G07 UC2-PrimColor 1675 549790 1789.N07.gz43 226512 F M0005638B:C07 UC2-PrimColor 1676 730660 1789.N07.gz43 226512 F M0005638G:A11 UC2-PrimColor 1676 730660 1789.N07.gz43 226512 F M0005638B:A11 UC2-PrimColor					M00056370B:C10	UC2-PrimColon
1665 487079 1789.F03.gz43 226440 F M00056374C:G08 UC2-PrimColon 1666 460284 1789.H03.gz43 226442 F M00056377A:C01 UC2-PrimColon 1667 725274 1789.103.gz43 226443 F M00056377A:C01 UC2-PrimColon 1668 553035 1789.N03.gz43 226448 F M0005637B:G01 UC2-PrimColon 1669 724009 1789.P03.gz43 226448 F M0005638B:B03 UC2-PrimColon 1670 568484 1789.B04.gz43 226452 F M0005638B:B03 UC2-PrimColon 1670 568484 1789.B04.gz43 226452 F M0005638B:F08 UC2-PrimColon 1671 725321 1789.L04.gz43 226462 F M0005638B:G08 UC2-PrimColon 1672 731542 1789.K05.gz43 226494 F M0005638B:G08 UC2-PrimColon 1673 725448 1789.D06.gz43 226494 F M0005638B:G08 UC2-PrimColon 1674 452806 1789.N07.gz43 226510 F M0005638B:G07 UC2-PrimColon 1675 549790 1789.L07.gz43 226510 F M0005638B:C17 UC2-PrimColon 1676 730660 1789.N07.gz43 226512 F M0005638B:A11 UC2-PrimColon 1676 730660 1789.N07.gz43 226512	$\overline{}$				M00056384D:A04	UC2-PrimColon
1666	_			_	M00056369B:E03	UC2-PrimColon
1667 725274 1789.103.gz43 226443 F M00056377D:F08 UC2-PrimColon				F	M00056374C:G08	UC2-PrimColon
1668 553035 1789.N03_gz43 226448 F M00056386A-G01 UC2-PrimColon 1669 724009 1789.P03_gz43 226450 F M0003638B-B03 UC2-PrimColon 1670 568484 1789.B04_gz43 226452 F M00056370B-E07 UC2-PrimColon 1671 725321 1789.L04_gz43 226462 F M0005638B-B03 UC2-PrimColon 1672 731542 1789.L05_gz43 226462 F M00056383B-G03 UC2-PrimColon 1673 725448 1789.L06_gz43 226494 F M00056383B-G03 UC2-PrimColon 1674 452806 1789.N06_gz43 226496 F M0005638B-G07 UC2-PrimColon 1675 549790 1789.L07_gz43 226510 F M0005638B-C07 UC2-PrimColon 1676 730660 1789.N07_gz43 226512 F M0005638B-C17 UC2-PrimColon 1676 730660 1789.N07_gz43 226512	1666	460284		F	M00056377A;C01	UC2-PrimColon
1669 724009 1789.P03.gz43 226450 F M00056338B:B03 UC2-PrimColon 1670 568484 1789.B04.gz43 226452 F M00056370B:E07 UC2-PrimColon 1671 725321 1789.L04.gz43 226462 F M00056338D:F08 UC2-PrimColon 1672 731542 1789.K05.gz43 226477 F M00056382C:F11 UC2-PrimColon 1673 725448 1789.L06.gz43 226494 F M00056383B:G08 UC2-PrimColon 1674 452806 1789.N06.gz43 226496 F M00056386E:A07 UC2-PrimColon 1675 549790 1789.L07.gz43 226512 F M0005638G:E07 UC2-PrimColon 1676 730660 1789.N07.gz43 226512 F M0005638G:A11 UC2-PrimColon 1789.N07.gz43 226512 F M000563	1667	725274	1789.I03.gz43_226443	F	M00056377D:F08	UC2-PrimColon
1670 568484 1789.B04_gz43 226452 F M00056370B:E07 UC2-PrimColon 1671 725321 1789.L04_gz43 226462 F M00056383B:F08 UC2-PrimColon 1672 731542 1789.K05_gz43 226477 F M00056383B:F08 UC2-PrimColon 1673 725448 1789.L06_gz43 226494 F M00056383B:G08 UC2-PrimColon 1674 452806 1789.N06_gz43 226496 F M0005638B:G07 UC2-PrimColon 1675 549790 1789.L07_gz43 226510 F M00056383C:E07 UC2-PrimColon 1676 730660 1789.N07_gz43 226512 F M0005638G:A11 UC2-PrimColon 1676 M0005638G:A11 UC2-PrimCol	1668	553035	1789.N03.gz43_226448	F	M00056386A:G01	UC2-PrimColon
1671 725321 1789.L04.gz43 226462 F M00056383B:F08 UC2-PrimColon 1672 731542 1789.K05.gz43 226477 F M00056382C:F11 UC2-PrimColon 1673 725448 1789.L06.gz43 226494 F M00056383B:G08 UC2-PrimColon 1674 452806 1789.N06.gz43 226494 F M00056386B:A07 UC2-PrimColon 1675 549790 1789.L07.gz43 226510 F M00056386B:A07 UC2-PrimColon 1676 730660 1789.N07.gz43 226512 F M00056386B:A11 UC2-PrimColon	1669	724009	1789.P03.gz43 226450	F	M00056388B;B03	UC2-PrimColon
1672 731542 1789.K05.gz43_226477 F M00056382C:F11 UC2-PrimColon 1673 725448 1789.L06.gz43_226494 F M00056383B:G08 UC2-PrimColon 1674 452806 1789.N06.gz43_226496 F M0005638B:A07 UC2-PrimColon 1675 549790 1789.L07.gz43_226510 F M0005638B:CE07 UC2-PrimColon 1676 730660 1789.N07.gz43_226512 F M0005638B:A11 UC2-PrimColon	1670	568484	1789.B04.gz43 226452	F	M00056370B:E07	UC2-PrimColon
1673 725448 1789.L06.gz43 226494 F M00056383B:G08 UC2-PrimColon 1674 452806 1789.N06.gz43 226496 F M00056386E:A07 UC2-PrimColon 1675 549790 1789.L07.gz43 226510 F M00056383C:E07 UC2-PrimColon 1676 730660 1789.N07.gz43 226512 F M0005638GE:A11 UC2-PrimColon	1671	725321	1789.L04.gz43 226462	F	M00056383B:F08	UC2-PrimColon
1673 725448 1789.L06.gz43 226494 F M00056383B:G08 UC2-PrimColon 1674 452806 1789.N06.gz43 226496 F M00056386E:A07 UC2-PrimColon 1675 549790 1789.L07.gz43 226510 F M00056383C:E07 UC2-PrimColon 1676 730660 1789.N07.gz43 226512 F M0005638GE:A11 UC2-PrimColon	1672	731542	1789.K05.gz43 226477	F	M00056382C:F11	UC2-PrimColon
1674 452806 1789.N06.gz43_226496 F M00056386B:A07 UC2-PrimColor 1675 549790 1789.L07.gz43_226510 F M00056383C:E07 UC2-PrimColor 1676 730660 1789.N07.gz43_226512 F M00056386B:A11 UC2-PrimColor	1673	725448		F	M00056383B:G08	
1675 549790 1789.L07.gz43 226510 F M00056383C:E07 UC2-PrimColon 1676 730660 1789.N07.gz43 226512 F M00056386B:A11 UC2-PrimColon	1674	452806	1789.N06.gz43 226496	F	M00056386B:A07	
1676 730660 1789.N07.gz43 226512 F M00056386B:A11 UC2-PrimColon	1675			F		UC2-PrimColon
	_					
1 10// 1 303623 1 1/89.B08.gz43 226516 1 F 1 M00056370C:F01 1 UC2-PrimColon	1677	503625	1789.B08.gz43 226516	F	M00056370C:F01	UC2-PrimColon
						UC2-PrimColon
						UC2-PrimColon
						UC2-PrimColon

Table 2

T	able 2				_
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ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
1681	727523	1789.M10.gz43_226559	F	M00056385A:A07	UC2-PrimColon
1682	459918	1789.N10.gz43_226560	F	M00056386B:D07	UC2-PrimColon
1683	735649	1789.B11.gz43_226564	F	M00056370D:C06	UC2-PrimColon
1684	734348	1789.F11.gz43_226568	F	M00056375A:D11	UC2-PrimColon
1685	558412	1789.O11.gz43_226577	F	M00056387C:D01	UC2-PrimColon
1686	724694	1789.H12.gz43_226586	F	M00056377C:A01	UC2-PrimColon
1687	723972	1789,J12.gz43_226588	F	M00056382A:A12	UC2-PrimColon
1688	623408	1789.N12.gz43_226592	F	M00056386B:E09	UC2-PrimColon
1689	640158	1790.F01.gz43_226792	F	M00056394C:D07	UC2-PrimColon
1690	725095	1790.J02.gz43_226812	F	M00056398C:D10	UC2-PrimColon
1691	562769	1790.O02.gz43_226817	F	M00056404D:G03	UC2-PrimColon
1692	479851	1790.L03.gz43_226830	F	M00056400C;G04	UC2-PrimColon
1693	481077	1790.N03.gz43_226832	F	M00056403C:F02	UC2-PrimColon
1694	656268	1790.K04.gz43_226845	F	M00056399D:C11	UC2-PrimColon
1695	724411	1790.N04.gz43_226848	F	M00056403C:F03	UC2-PrimColon
1696	640261	1790.N05.gz43_226864	F	M00056403C:F12	UC2-PrimColon
1697	426297	1790.D07.gz43_226886	F	M00056392D:C05	UC2-PrimColon
1698	649591	1790.M07.gz43_226895	F	M00056402C:B09	UC2-PrimColon
1699	729111	1790.D08.gz43_226902	F	M00056392D:H02	UC2-PrimColon
1700	733563	1790.G08.gz43_226905	F	M00056396A:D03	UC2-PrimColon
1701	729637	1790.A10.gz43_226931	F	M00056389B:H03	UC2-PrimColon
1702	726494	1790,K10.gz43_226941	F	M00056400A:D08	UC2-PrimColon
1703	724714	1790.M10.gz43_226943	F	M00056402D:A01	UC2-PrimColon
1704	543323	1790.P10.gz43_226946	F	M00056406C:D09	UC2-PrimColon
1705	556769	1790.B11.gz43_226948	F	M00056390C:B08	UC2-PrimColon
1706	209	1790.M11.gz43_226959	F	M00056402D:B07	UC2-PrimColon
1707	724914	1790.B12.gz43_226964	F	M00056390C:C04	UC2-PrimColon
1708	614463	1790.D12.gz43_226966	F	M00056393A:G06	UC2-PrimColon
1709	474981	1790.G12.gz43_226969	F	M00056396A:G10	UC2-PrimColon
1710	729950	1790.L12.gz43_226974	F	M00056401A:F09	UC2-PrimColon
1711	640756	1790.N12.gz43_226976	F	M00056404A:G03	UC2-PrimColon
1712	449228	1790.O13.gz43_226993	F	M00056405C:H04	UC2-PrimColon
1713	423420	1790.P13.gz43_226994	F	M00056406C:F12	UC2-PrimColon
1714	727150	1790.C14.gz43_226997	F	M00056391C:H12	UC2-PrimColon
1715	503173	1790.E14.gz43_226999	F	M00056394B:C11	UC2-PrimColon
1716	188309	1790.N14.gz43_227008	F	M00056404B:B06	UC2-PrimColon
1717	448758	1790.J15.gz43_227020	F	M00056399B:G09	UC2-PrimColon
1718	735234	1790.L15.gz43_227022	F	M00056401B:B01	UC2-PrimColon
1719	483147	1790.P15.gz43_227026	F	M00056407A:D10	UC2-PrimColon
1720	642985	1790.G16.gz43_227033	F	M00056396B:E10	UC2-PrimColon
1721	725451	1790.K16.gz43_227037	F	M00056400A:G10	UC2-PrimColon
1722	729809	1790.P16.gz43 227042	F	M00056407A:G03	UC2-PrimColon

Table 2

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1723	495712	1790.A17.gz43 227043	F	M00056389D:F01	UC2-PrimColon
1724	730948	1790.O17.gz43 227057	F	M00056405D:E05	UC2-PrimColon
1725	457508	1790.C18.gz43 227061	F	M00056391D:E06	UC2-PrimColon
1726	417549	1790.M18.gz43_227071	F	M00056403A:E09	UC2-PrimColon
1727	725454	1790.E19.gz43_227079	F	M00056394B:G05	UC2-PrimColon
1728	448817	1790.C20.gz43_227093	F	M00056392A:F06	UC2-PrimColon
1729	726411	1790.O20.gz43 227105	F	M00056406A:C02	UC2-PrimColon
1730	558794	1790.D21.gz43_227110	F	M00056393D:A09	UC2-PrimColon
1731	641687	1790.P21.gz43_227122	F	M00056409A:C03	UC2-PrimColon
1732	733643	1790.N22.gz43_227136	F	M00056404C:C12	UC2-PrimColon
1733	523364	1790.F23.gz43_227144	F	M00056395C:H03	UC2-PrimColon
1734	468467	1790.M23.gz43_227151	F	M00056403B:D09	UC2-PrimColon
1735	402799	1790.O23.gz43_227153	F	M00056406A:E01	UC2-PrimColon
1736	470462	1790.P23.gz43_227154	F	M00056409A:D03	UC2-PrimColon
1737	118514	1790.D24.gz43_227158	F	M00056393D:C12	UC2-PrimColon
1738	732821	1790.F24.gz43_227160	F	M00056395D:B01	UC2-PrimColon
1739	447222	1790.K24.gz43_227165	F	M00056400C:B11	UC2-PrimColon
1740	729779	1790.M24.gz43_227167	F	M00056403B:G12	UC2-PrimColon
1741	727633	1790.P24.gz43_227170	F	M00056409A:D06	UC2-PrimColon
1742	475797	1791.F01.gz43_227176	F	M00056415A:D03	UC2-PrimColon
1743	648170	1791.G01.gz43_227177	F	M00056416B;B11	UC2-PrimColon
1744	472684	1791.I01.gz43_227179	F	M00056418A:G09	UC2-PrimColon
1745	477718	1791.O02.gz43_227201	F	M00056423D:H07	UC2-PrimColon
1746	729784	1791.A03.gz43_227203	F	M00056409B:C12	UC2-PrimColon
1747	402471	1791.F03.gz43_227208	F	M00056415B:B10	UC2-PrimColon
1748	453762	1791.K03.gz43_227213	F	M00056419D:G04	UC2-PrimColon
1749	511149	1791.M03.gz43_227215	F	M00056422B:A08	UC2-PrimColon
1750	727090	1791.G04.gz43_227225	F	M00056416B:H01	UC2-PrimColon
1751	560957	1791.K04.gz43_227229	F	M00056420A:C01	UC2-PrimColon
1752	727845	1791.M04.gz43_227231	F	M00056422B:C12	UC2-PrimColon
1753	642170	1791.H05.gz43_227242	F	M00056417B:D05	UC2-PrimColon
1754	128749	1791.K05.gz43_227245	F	M00056420A:C12	UC2-PrimColon
1755	736632	1791.P05.gz43_227250	F	M00056424D:C08	UC2-PrimColon
1756	542957	1791.J07.gz43_227276	F	M00056419A:F08	UC2-PrimColon
1757	733490	1791.008.gz43_227297	F	M00056424A;F05	UC2-PrimColon
1758	649722	1791.009.gz43_227313	F	M00056424A:F12	UC2-PrimColon
1759	562137	1791.L11.gz43_227342	F	M00056421C:E06	UC2-PrimColon
1760	729199	1791.G12.gz43_227353	F	M00056416D;C08	UC2-PrimColon
1761	730178	1791.L12.gz43_227358	F	M00056421C;E12	UC2-PrimColon
1762	435349	1791.G13.gz43_227369	F	M00056416D:D12	UC2-PrimColon
1763	723963	1791.P13.gz43_227378	F	M00056425B:B02	UC2-PrimColon
1764	725408	1791.H14.gz43_227386	F	M00056417C:G10	UC2-PrimColon

Table 2

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SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
1765	500253	1791.L14.gz43_227390	F	M00056421C:H01	UC2-PrimColon
1766	644723	1791.N14.gz43_227392	F	M00056423C:G05	UC2-PrimColon
1767	474965	1791.M15.gz43_227407	F	M00056422C:B10	UC2-PrimColon
1768	724024	1791.N17.gz43_227440	F	M00056423D:B06	UC2-PrimColon
1769	27083	1791.D18.gz43_227446	F	M00056413B:D07	UC2-PrimColon
1770	729899	1791.E18.gz43_227447	F	M00056414C:H08	UC2-PrimColon
1771	728986	1791.G18.gz43_227449	F	M00056416D:H10	UC2-PrimColon
1772	640744	1791.A19.gz43_227459	F	M00056410A:A04	UC2-PrimColon
1773	703978	1791.B19.gz43_227460	F	M00056411B:D01	UC2-PrimColon
1774	725408	1791.O19.gz43_227473	F	M00056424C:B10	UC2-PrimColon
1775	725407	1791.H20.gz43_227482	F	M00056417D:G09	UC2-PrimColon
1776	736507	1791.K21.gz43_227501	F	M00056420C:E08	UC2-PrimColon
1777	555359	1791.O21.gz43_227505	F	M00056424C:F02	UC2-PrimColon
1778	724773	1791.C22.gz43_227509	F	M00056412B:B08	UC2-PrimColon
1779	724220	1791.P22.gz43_227522	F	M00056425D:D08	UC2-PrimColon
1780	641383	1791.A23.gz43_227523	F	M00056410B:A09	UC2-PrimColon
1781	562581	1791.K23.gz43_227533	F	M00056420D:A09	UC2-PrimColon
1782	726768	1791.P23.gz43_227538	F	M00056425D:H01	UC2-PrimColon
1783	729579	1791.H24.gz43_227546	F	M00056418A:G01	UC2-PrimColon
1784	498194	1791.L24.gz43_227550	F	M00056422A:F07	UC2-PrimColon
1785	288134	1791.N24.gz43_227552	F	M00056423D:F10	UC2-PrimColon
1786	553983	1792.P01.gz43_227582	F	M00056486B:G02	UC2-PrimColon
1787	724400	1792.E03.gz43_227603	F	M00056433B:A09	UC2-PrimColon
1788	645476	1792.J05.gz43_227640	F	M00056479C:E04	UC2-PrimColon
1789	556115	1792.D06.gz43_227650	F	M00056432A:A01	UC2-PrimColon
1790	734287	1792.F06.gz43_227652	F	M00056434C:B01	UC2-PrimColon
1791	725584	1792.I07.gz43_227671	F	M00056478C:H01	UC2-PrimColon
1792	725006	1792.P07.gz43_227678	F	M00056486D:A12	UC2-PrimColon
1793	731158	1792.A09.gz43_227695	F	M00056426B:G12	UC2-PrimColon
1794	592186	1792.B09.gz43_227696	F	M00056428A:F10	UC2-PrimColon
1795	452618	1792.M09.gz43_227707	F	M00056483A:F10	UC2-PrimColon
1796	725238	1792.G10.gz43_227717	F	M00056476C:E05	UC2-PrimColon
1797	727948	1792.P10.gz43_227726	F	M00056486D:F04	UC2-PrimColon
1798	452981	1792.A11.gz43_227727	F	M00056426C:G05	UC2-PrimColon
1799	710362	1792.B11.gz43_227728	F	M00056428B:B01	UC2-PrimColon
1800	480005	1792.C11.gz43_227729	F	M00056429D:E02	UC2-PrimColon
1801	471931	1792.P11.gz43_227742	F	M00056487A:C10	UC2-PrimColon
1802	725454	1792.H12.gz43_227750	F	M00056477C:G09	UC2-PrimColon
1803	729584	1792.I12.gz43_227751	F	M00056478D:G02	UC2-PrimColon
1804	725210	1792.N12.gz43_227756	F	M00056484B:E11	UC2-PrimColon
1805	462687	1792.P12.gz43_227758	F	M00056487A:D01	UC2-PrimColon
1806	724183	1792.M13.gz43_227771	F	M00056483B:E04	UC2-PrimColon

Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
1807	724390	1792.M14.gz43_227787	F	M00056483B:F04	UC2-PrimColon
1808	647375	1792.G16.gz43_227813	F	M00056476D:H11	UC2-PrimColon
1809	736751	1792.G17.gz43_227829	F	M00056477A:B11	UC2-PrimColon
1810	725348	1792.P17.gz43_227838	F	M00056487A:F05	UC2-PrimColon
1811	481594	1792.L18.gz43_227850	F	M00056482B:B11	UC2-PrimColon
1812	675299	1792.N18.gz43_227852	F	M00056484D:D01	UC2-PrimColon
1813	734646	1792.K19.gz43_227865	F	M00056481A:F02	UC2-PrimColon
1814	453713	1792.L20.gz43_227882	F	M00056482B:C09	UC2-PrimColon
1815	724183	1792.M20.gz43_227883	F	M00056483C:D06	UC2-PrimColon
1816	730282	1792.M24.gz43_227947	F	M00056483D:F06	UC2-PrimColon
1817	733006	1801.D01.gz43_227954	F	M00056491B:B09	UC2-PrimColon
1818	733151	1801.I01.gz43_227959	F	M00056496D:B12	UC2-PrimColon
1819	734261	1801.J01.gz43_227960	F	M00056497C:D05	UC2-PrimColon
1820	514142	1801.K01.gz43_227961	F	M00056498D:C01	UC2-PrimColon
1821	471232	1801.M01.gz43_227963	F	M00056501B:B09	UC2-PrimColon
1822	447785	1801.D02.gz43_227970	F	M00056491B;E02	UC2-PrimColon
1823	523753	1801.J02.gz43_227976	F	M00056497C:E01	UC2-PrimColon
1824	451383	1801.M02.gz43_227979	F	M00056501B:C07	UC2-PrimColon
1825	635951	1801.K03.gz43_227993	F	M00056499A:A04	UC2-PrimColon
1826	731820	1801.A04.gz43_227999	F	M00056487C:E03	UC2-PrimColon
1827	446928	1801.E04.gz43_228003	F	M00056492B;D06	UC2-PrimColon
1828	732598	1801.P04.gz43_228014	F	M00056504B;B01	UC2-PrimColon
1829	725434	1801.B05.gz43_228016	F	M00056488B:G10	UC2-PrimColon
1830	460690	1801.G05.gz43_228021	F	M00056495A:C02	UC2-PrimColon
1831	611604	1801.J05.gz43_228024	F	M00056497D:C11	UC2-PrimColon
1832	635951	1801.L05.gz43_228026	F	M00056500A:G12	UC2-PrimColon
1833	429191	1801.B06.gz43_228032	F	M00056488B;H04	UC2-PrimColon
1834	640116	1801.O06.gz43_228045	F	M00056503B:G11	UC2-PrimColon
1835	734043	1813.H05.gz43_229558	F	M00056576A:A04	UC2-PrimColon
1836	508210	1813.I06.gz43_229575	F	M00056577A;F10	UC2-PrimColon
1837	11632	1813.O07.gz43_229597	F	M00056585C:C08	UC2-PrimColon
1838	475757	1813.H08.gz43_229606	F	M00056576A;E01	UC2-PrimColon
1839	729281	1813.J10.gz43_229640	F	M00056578D:A02	UC2-PrimColon
1840	733970	1813.N10.gz43_229644	F	M00056584C:A06	UC2-PrimColon
1841	727410	1813.A13.gz43_229679	F	M00056567A:H06	UC2-PrimColon
1842	465446	1813.L13.gz43_229690	F	M00056581D:A08	UC2-PrimColon
1843	511351	1813.M13.gz43_229691	F	M00056583C:A05	UC2-PrimColon
1844	733664	1813.I20.gz43_229799	F	M00056577D:F08	UC2-PrimColon
1845	548893	1813.C22.gz43_229825	F	M00056570A:D09	UC2-PrimColon
1846	513540	1813.H22.gz43 229830	F	M00056576C:G01	UC2-PrimColon
1847	725759	1813.I22.gz43_229831	F	M00056578A:B05	UC2-PrimColon
1848	726104	1816.C19.gz43_230929	F	M00056633D:E05	UC2-PrimColon

Table 2

1	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
1849	482868	1816.E19.gz43 230931	F	M00056636C:H05	UC2-PrimColon
1850	727332	1816.I19.gz43_230935	F	M00056641C:G09	UC2-PrimColon
1851	727078	1816.K19.gz43_230937	F	M00056644C:B03	UC2-PrimColon
1852	727108	1816.L19.gz43_230938	F	M00056645D:E06	UC2-PrimColon
1853	736325	1816.B20.gz43_230944	F	M00056632C;E03	UC2-PrimColon
1854	726818	1816.J20.gz43_230952	F	M00056642D:C05	UC2-PrimColon
1855	724083	1816.M20.gz43_230955	F	M00056646D:E03	UC2-PrimColon
1856	724565	1816.N20.gz43 230956	F	M00056648B:C08	UC2-PrimColon
1857	207530	1816.O20.gz43_230957	F	M00056650A:E01	UC2-PrimColon
1858	736778	1816.C21.gz43_230961	F	M00056634A:C06	UC2-PrimColon
1859	559684	1816.E21.gz43_230963	F	M00056636D:F03	UC2-PrimColon
1860	726494	1816.B22.gz43_230976	F	M00056632C:H08	UC2-PrimColon
1861	419711	1816.G22.gz43_230981	F	M00056639A:F11	UC2-PrimColon
1862	377935	1816.M22.gz43 230987	F	M00056646D:F07	UC2-PrimColon
1863	726448	1816.C23.gz43 230993	F	M00056634A:G11	UC2-PrimColon
1864	215249	1816.G23.gz43 230997	F	M00056639A:H12	UC2-PrimColon
1865	735322	1816.N23.gz43 231004	F	M00056648B:E09	UC2-PrimColon
1866	727175	1816.O23.gz43 231005	F	M00056650B:C11	UC2-PrimColon
1867	731966	1816.P23.gz43 231006	F	M00056651B:E12	UC2-PrimColon
1868	727944	1816.B24.gz43 231008	F	M00056632D:E02	UC2-PrimColon
1869	730596	1816.M24.gz43 231019	F	M00056647A:F06	UC2-PrimColon
1870	725607	1825.N13.gz43_231228	F	M00056667A:A12	UC2-PrimColon
1871	45934	1825.P13.gz43 231230	F	M00056670A;G02	UC2-PrimColon
1872	484987	1825.F14.gz43 231236	F	M00056658A:E11	UC2-PrimColon
1873	481592	1825.M14.gz43 231243	F	M00056666A;C04	UC2-PrimColon
1874	735605	1825.O14.gz43 231245	F	M00056668D:E03	UC2-PrimColon
1875	734184	1825.P14.gz43 231246	F	M00056670B:A12	UC2-PrimColon
1876	477718	1825.E16.gz43 231267	F	M00056657A:F06	UC2-PrimColon
1877	552927	1825.F16.gz43_231268	F	M00056658B:C03	UC2-PrimColon
1878	719620	1825.J16.gz43_231272	F	M00056663A:B02	UC2-PrimColon
1879	27083	1825.M16.gz43 231275	F	M00056666A;D09	UC2-PrimColon
1880	478458	1825.P17.gz43 231294	F	M00056670B:G05	UC2-PrimColon
1881	448104	1825.L19.gz43_231322	F	M00056664D;H06	UC2-PrimColon
1882	729675	1825.J20.gz43 231336	F	M00056663A:H09	UC2-PrimColon
1883	730570	1825.B21.gz43 231344	F	M00056653D:F07	UC2-PrimColon
1884	727108	1825.D21.gz43 231346	F	M00056656A;E01	UC2-PrimColon
1885	447074	1825,I21.gz43 231351	F	M00056661D:D06	UC2-PrimColon
1886	463741	1825.F22.gz43 231364	F	M00056658C:B12	UC2-PrimColon
1887	732351	1825.N22.gz43 231372	F	M00056667D:E05	UC2-PrimColon
1888	727343	1825,L23.gz43 231386	F	M00056665A:G08	UC2-PrimColon
1889	736415	1825,B24.gz43_231392	F	M00056654A:A07	UC2-PrimColon
1890	554597	1825.O24.gz43 231405	F	M00056669B:E10	UC2-PrimColon

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Table 2

T	able 2				
SEQ			ORIEN		
ш ио	CLUSTER	SEQ NAME	Т	CLONE ID	LIBRARY
1891	723914	1826.A13.gz43_231599	_ F	M00056672A:E05	UC2-PrimColon
1892	734622	1826.C14.gz43_231617	F	M00056674D:E09	UC2-PrimColon
1893	645925	1826.I14.gz43_231623	F	M00056683B:H04	UC2-PrimColon
1894	641777	1826.D16.gz43 231650	F	M00056676B:G12	UC2-PrimColon
1895	734828	1826.E17.gz43_231667	F	M00056678B:E11	UC2-PrimColon
1896	381623	1826.I17.gz43_231671	F	M00056683C:H10	UC2-PrimColon
1897	517274	1826.J17.gz43_231672	F	M00056685A:H11	UC2-PrimColon
1898	446575	1826.C18.gz43_231681	F	M00056675A:C02	UC2-PrimColon
1899	725899	1826.D18.gz43_231682	F	M00056676C:B04	UC2-PrimColon
1900	492094	1826.M19.gz43_231707	F	M00056689B:F03	UC2-PrimColon
1901	736246	1826.E20.gz43_231715	F	M00056678B:H02	UC2-PrimColon
1902	473238	1826.J20.gz43_231720	F	M00056685B:G04	UC2-PrimColon
1903	730829	1826.K20.gz43_231721	F	M00056686D:E04	UC2-PrimColon
1904	726722	1826.A21.gz43_231727	F	M00056672D:B08	UC2-PrimColon
1905	492629	1826.021.gz43_231741	F	M00056692A:A05	UC2-PrimColon
1906	730664	1826.E22.gz43_231747	F	M00056678C:A12	UC2-PrimColon
1907	729387	1826.G22.gz43_231749	F	M00056681A:E06	UC2-PrimColon
1908	546121	1826.A23.gz43_231759	F	M00056672D:E04	UC2-PrimColon
1909	513156	1826.C23.gz43_231761	F	M00056675A:G02	UC2-PrimColon
1910	728791	1826.G24.gz43_231781	F	M00056681B;A11	UC2-PrimColon
1911	559610	1826.I24.gz43_231783	F	M00056684B:B12	UC2-PrimColon
1912	729205	1826.J24.gz43_231784	F	M00056685C:G07	UC2-PrimColon
1913	561301	1827.C01.gz43_231793	F	M00056695C:C04	UC2-PrimColon
1914	728556	1827.E01.gz43_231795	F	M00056697C:H10	UC2-PrimColon
1915	640458	1827.I01.gz43_231799	F	M00056701C:G09	UC2-PrimColon
1916	650136	1827.K01.gz43_231801	F	M00056703C:F04	UC2-PrimColon
1917	288801	1827.L01.gz43_231802	F	M00056704D:H10	UC2-PrimColon
1918	648777	1827.M01.gz43_231803	F	M00056706A:B05	UC2-PrimColon
1919	728196	1827.G02.gz43_231813	F	M00056699C:F09	UC2-PrimColon
1920	644354	1827.I02.gz43_231815	F	M00056701D:C03	UC2-PrimColon
1921	419465	1827.M02.gz43_231819	F	M00056706A:E09	UC2-PrimColon
1922	558317	1827.P02.gz43_231822	F	M00056708C:F11	UC2-PrimColon
1923	693869	1827.B03.gz43_231824	F	M00056694A:G03	UC2-PrimColon
1924	553805	1827.L03.gz43_231834	F	M00056705A:B12	UC2-PrimColon
1925	626791	1827.N03.gz43_231836	F	M00056707A:G11	UC2-PrimColon
1926	580601	1827.A04.gz43_231839	F	M00056693B:D11	UC2-PrimColon
1927	493830	1827.B04.gz43_231840	F	M00056694A:G11	UC2-PrimColon
1928	724714	1827.E04.gz43_231843	F	M00056697D:B09	UC2-PrimColon
1929	737109	1827.F04.gz43_231844	F	M00056698D:E11	UC2-PrimColon
1930	395604	1827.G04.gz43_231845	F	M00056699C:G11	UC2-PrimColon
1931	735834	1827.J04.gz43_231848	F	M00056702D:G03	UC2-PrimColon
1932	726852	1827.L04.gz43_231850	F	M00056705A:D02	UC2-PrimColon

Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	Т	CLONE ID	LIBRARY
1933	728892	1827,P05.gz43 231870	F	M00056708D:B03	UC2-PrimColon
1934	540618	1827.C06.gz43 231873	F	M00056695C:H08	UC2-PrimColon
1935	501030	1827.E06.gz43_231875	F	M00056697D:C09	UC2-PrimColon
1936	477366	1827.F06.gz43 231876	F	M00056698D:G09	UC2-PrimColon
1937	727948	1827.I06.gz43 231879	F	M00056701D:D04	UC2-PrimColon
1938	649927	1827.K06.gz43_231881	F	M00056703D:F08	UC2-PrimColon
1939	449752	1827,M06.gz43_231883	F	M00056706B;C02	UC2-PrimColon
1940	649106	1827,N06,gz43_231884	F	M00056707B:E02	UC2-PrimColon
1941	730463	1827.P06.gz43_231886	F	M00056708D:D10	UC2-PrimColon
1942	462687	1827.E07.gz43_231891	F	M00056697D:C12	UC2-PrimColon
1943	553140	1827.I07.gz43_231895	F	M00056701D:G03	UC2-PrimColon
1944	736100	1827.M07.gz43_231899	F	M00056706B;C07	UC2-PrimColon
1945	730375	1827.B08.gz43_231904	F	M00056694B:H10	UC2-PrimColon
1946	403671	1827.K08.gz43_231913	F	M00056704A:B05	UC2-PrimColon
1947	595181	1827.M08.gz43_231915	F	M00056706B:D04	UC2-PrimColon
1948	573733	1827.A09.gz43_231919	F	M00056693B:H03	UC2-PrimColon
1949	550454	1827.I09.gz43_231927	F	M00056701D:H12	UC2-PrimColon
1950	651050	1827.J09.gz43_231928	F	M00056703A:G01	UC2-PrimColon
1951	728768	1827.L09.gz43_231930	F	M00056705B:A12	UC2-PrimColon
1952	736276	1827.N09.gz43_231932	F	M00056707B:F06	UC2-PrimColon
1953	472837	1827.C10.gz43_231937	F	M00056695D:E05	UC2-PrimColon
1954	732770	1827.H10.gz43_231942	F	M00056700D:H07	UC2-PrimColon
1955	83388	1827.L10.gz43_231946	F	M00056705B;D10	UC2-PrimColon
1956	584693	1827.M10.gz43_231947	F	M00056706C:A07	UC2-PrimColon
1957	588059	1827.D11.gz43_231954	F	M00056697A:D02	UC2-PrimColon
1958	452662	1827.F11.gz43_231956	F	M00056699A;D08	UC2-PrimColon
1959	728898	1827.M11.gz43_231963	F	M00056706C:B12	UC2-PrimColon
1960	672032	1827.F12.gz43_231972	F	M00056699A:E08	UC2-PrimColon
1961	496084	1827.H12.gz43_231974	F	M00056701A;A02	UC2-PrimColon
1962	736579	1827.O12.gz43_231981	F	M00056708B;B09	UC2-PrimColon
1963	727968	1827.A13.gz43_231983	F	M00056693C:D12	UC2-PrimColon
1964	447624	1827,J13.gz43_231992	F	M00056703B:A04	UC2-PrimColon
1965	728552	1827.K13.gz43_231993	F	M00056704A:H08	UC2-PrimColon
1966	733064	1827.M13.gz43_231995	F	M00056706C:D05	UC2-PrimColon
1967	607202	1827.O13.gz43_231997	F	M00056708B;D03	UC2-PrimColon
1968	473433	1827.P13.gz43_231998	F	M00056708D:H06	UC2-PrimColon
1969	617813	1827.A14.gz43_231999	F	M00056693C:G06	UC2-PrimColon
1970	728300	1827.G14.gz43_232005	F	M00056700A:F12	UC2-PrimColon
1971	727760	1827.K14.gz43_232009	F	M00056704B:C02	UC2-PrimColon
1972	454087	1827.P14.gz43_232014	F	M00056709A:A05	UC2-PrimColon
1973	425396	1827.A15.gz43_232015	F	M00056693D:B02	UC2-PrimColon
1974	552430	1827.N15.gz43_232028	F	M00056707C:E01	UC2-PrimColon

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T	able 2				
CEO.					
SEQ			ORIEN		
	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
1975	650542	1827.C16.gz43_232033	F	M00056696A:G12	UC2-PrimColon
1976	725084	1827.D16.gz43_232034	F	M00056697A:H10	UC2-PrimColon
1977	446607	1827.G16.gz43_232037	F	M00056700A:G05	UC2-PrimColon
1978	4584	1827.H16.gz43_232038	F	M00056701A:G09	UC2-PrimColon
1979	560678	1827.J16.gz43_232040	F	M00056703B:D07	UC2-PrimColon
1980	730639	1827.N16.gz43_232044	F	M00056707C:E03	UC2-PrimColon
1981	727506	1827.C17.gz43_232049	F	M00056696B:B03	UC2-PrimColon
1982	540618	1827.E17.gz43_232051	F	M00056698B:E10	UC2-PrimColon
1983	735676	1827.I17.gz43_232055	F	M00056702C:A10	UC2-PrimColon
1984	728768	1827.N17.gz43_232060	F	M00056707C:F10	UC2-PrimColon
1985	475942	1827.A18.gz43_232063	F	M00056693D:E12	UC2-PrimColon
1986	631056	1827.H18.gz43_232070	F	M00056701B:D02	UC2-PrimColon
1987	259218	1827.P18.gz43_232078	F	M00056709A:H11	UC2-PrimColon
1988	724489	1827.H19.gz43_232086	F	M00056701B:D06	UC2-PrimColon
1989	727845	1827.D21.gz43_232114	F	M00056697C:D11	UC2-PrimColon
1990	641072	1827.F21.gz43_232116	F	M00056699B:G04	UC2-PrimColon
1991	553483	1827.A22.gz43_232127	F	M00056694A:C05	UC2-PrimColon
1992	551096	1827.C24.gz43_232161	F	M00056696B:H01	UC2-PrimColon
1993	727649	1827.F24.gz43_232164	F	M00056699C:C02	UC2-PrimColon
1994	728196	1827.H24.gz43_232166	F	M00056701C:F08	UC2-PrimColon
1995	729428	1828.A01.gz43_232175	F	M00056709C:F06	UC2-PrimColon
1996	724050	1828.F01.gz43_232180	F	M00056714B:C12	UC2-PrimColon
1997	732351	1828.L01.gz43_232186	F	M00056721C:E05	UC2-PrimColon
1998	730627	1828.M01.gz43_232187	F	M00056722C:C09	UC2-PrimColon
1999	728182	1828.H02.gz43_232198	F	M00056716B:F12	UC2-PrimColon
2000	727878	1828.M02.gz43_232203	F	M00056722C:D11	UC2-PrimColon
2001	462687	1828.N02.gz43_232204	F	M00056723C:C09	UC2-PrimColon
2002	555763	1828.L03.gz43_232218	F	M00056721C:H01	UC2-PrimColon
2003	728627	1828.P03.gz43_232222	F	M00056725C:H06	UC2-PrimColon
2004	630516	1828.H04.gz43_232230	F	M00056716C:B06	UC2-PrimColon
2005	584179	1828.A05.gz43_232239	F	M00056709D:D05	UC2-PrimColon
2006	558544	1828.G05.gz43_232245	F	M00056715C:B07	UC2-PrimColon
2007	553294	1828.N05.gz43_232252	F	M00056723C:E01	UC2-PrimColon
2008	726576	1828.O05.gz43_232253	F	M00056724C:H11	UC2-PrimColon
2009	734622	1828.L07.gz43_232282	F	M00056721D:D01	UC2-PrimColon
2010	735362	1828,P07.gz43_232286	F	M00056726A:C12	UC2-PrimColon
2011	455297	1828.A08.gz43_232287	F	M00056709D:E12	UC2-PrimColon
2012	734990	1828.N08.gz43_232300	F	M00056723C:G03	UC2-PrimColon
2013	707609	1828.B10.gz43_232320	F	M00056710D:C05	UC2-PrimColon
2014	728178	1828.L10.gz43_232330	F	M00056721D:F12	UC2-PrimColon
2015	447634	1828.P10.gz43_232334	F	M00056726B:H06	UC2-PrimColon
2016	729813	1828.F11.gz43_232340	F	M00056714D:A11	UC2-PrimColon

Table 2

SEQ			·		l
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ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
2017	728528	1828.I11.gz43 232343	F	M00056718A:H05	UC2-PrimColon
2018	732770	1828,J11.gz43 232344	F	M00056719B:G04	UC2-PrimColon
2019	505858	1828.F12.gz43 232356	F	M00056714D:E08	UC2-PrimColon
2020	735047	1828.E13.gz43 232371	F	M00056714A:B02	UC2-PrimColon
2021	727749	1828.K13.gz43 232377	F	M00056721A:C07	UC2-PrimColon
2022	598746	1828.L13.gz43 232378	F	M00056722A;E10	UC2-PrimColon
2023	728061	1828.O13.gz43 232381	F	M00056725A:E02	UC2-PrimColon
2024	448217	1828.F14.gz43 232388	F	M00056714D:H07	UC2-PrimColon
2025	485899	1828.L14.gz43_232394	F	M00056722A:F08	UC2-PrimColon
2026	641287	1828.B15.gz43 232400	F	M00056711A:A09	UC2-PrimColon
2027	48619	1828.D15.gz43 232402	F	M00056713A:F05	UC2-PrimColon
2028	728929	1828.E15.gz43 232403	F	M00056714A:B09	UC2-PrimColon
2029	451972	1828.I15.gz43 232407	F.	M00056718B:C02	UC2-PrimColon
2030	486363	1828.J18.gz43 232456	F	M00056719C:B10	UC2-PrimColon
2031	728897	1828.C19.gz43 232465	F	M00056712B;B11	UC2-PrimColon
2032	448217	1828.E19.gz43 232467	F	M00056714A:E12	UC2-PrimColon
2033	646309	1828.I19.gz43 232471	F	M00056718C:B01	UC2-PrimColon
2034	728303	1828.J19.gz43 232472	F	M00056719C:F06	UC2-PrimColon
2035	728002	1828.K19.gz43 232473	F	M00056721B:D03	UC2-PrimColon
2036	473640	1828.N20.gz43 232492	F	M00056724B:E11	UC2-PrimColon
2037	727480	1828.O21.gz43 232509	F	M00056725C:A03	UC2-PrimColon
2038	509678	1828.P21.gz43_232510	F	M00056726D:G08	UC2-PrimColon
2039	728445	1828.E22.gz43 232515	F	M00056714A:H06	UC2-PrimColon
2040	447150	1828.N22.gz43_232524	F	M00056724B:G03	UC2-PrimColon
2041	728273	1828.A23.gz43_232527	F	M00056710B;F05	UC2-PrimColon
2042	734453	1828.C23.gz43_232529	F	M00056712B:F02	UC2-PrimColon
2043	732006	1828.D23.gz43_232530	F	M00056713C:A10	UC2-PrimColon
2044	449585	1828.I23.gz43_232535	F	M00056718D:D12	UC2-PrimColon
2045	728966	1828.E24.gz43_232547	F	M00056714B:B06	UC2-PrimColon
2046	529356	1838.C01.gz43_232945	F	M00056747D:A03	UC2-PrimColon
2047	551693	1838.P01.gz43_232958	F	M00056764C:A02	UC2-PrimColon
2048	438663	1838.B03.gz43_232976	F	M00056746D:D06	UC2-PrimColon
2049	735464	1838.E03.gz43_232979	F	M00056750B:H03	UC2-PrimColon
2050	736035	1838.J03.gz43_232984	F	M00056756C:D06	UC2-PrimColon
2051	736210	1838.L03.gz43_232986	F	M00056759C:C04	UC2-PrimColon
2052	728464	1838.I05.gz43_233015	F	M00056755B:E07	UC2-PrimColon
2053	730178	1838.K05.gz43_233017	F	M00056758C:B08	UC2-PrimColon
2054	481614	1838,N05.gz43_233020	F	M00056762B:D06	UC2-PrimColon
2055	402799	1839.A01.gz43_233327	F	M00056766D:G09	UC2-PrimColon
2056	675768	1839.N01.gz43_233340	F	M00056784D:C06	UC2-PrimColon
2057	577305	1839.P01.gz43_233342	F	M00056787B:C07	UC2-PrimColon
2058	730505	1839.K02.gz43_233353	F	M00056781A:E06	UC2-PrimColon

Table 2

Table 2					
SEQ			ORIEN		
ID NO	0-001-20	SEQ NAME	T	CLONE ID	LIBRARY
2059	349977	1839.B03.gz43_233360	F	M00056768D:A09	UC2-PrimColon
2060	726344	1839.N03.gz43_233372	F	M00056784D:G10	UC2-PrimColon
2061	724183	1839.J04.gz43_233384	F	M00056780B:E06	UC2-PrimColon
2062	730352	1839.M04.gz43_233387	F	M00056783C:E03	UC2-PrimColon
2063	510596	1839.E05.gz43_233395	F	M00056773C:C09	UC2-PrimColon
2064	128773	1839.F05.gz43_233396	F	M00056775A:A05	UC2-PrimColon
2065	728464	1839.G06.gz43_233413	F	M00056776A:G04	UC2-PrimColon
2066	449010	1839.L06.gz43_233418	F	M00056782B:G05	UC2-PrimColon
2067	460023	1839.M06.gz43_233419	F	M00056783D:A02	UC2-PrimColon
2068	733744	1839.K07.gz43_233433	F	M00056781B:C03	UC2-PrimColon
2069	2180	1839.L07.gz43_233434	F	M00056782B:H11	UC2-PrimColon
2070	556458	1839,M07.gz43_233435	F	M00056783D:B07	UC2-PrimColon
2071	711325	1839.A08.gz43_233439	F	M00056767A:F12	UC2-PrimColon
2072	135593	1839.E08.gz43_233443	F	M00056773C:F12	UC2-PrimColon
2073	639178	1839.K08.gz43_233449	F	M00056781B:C05	UC2-PrimColon
2074	504513	1839.B09.gz43_233456	F	M00056769C:C03	UC2-PrimColon
2075	493193	1839.J09.gz43_233464	F	M00056780B:H04	UC2-PrimColon
2076	728133	1839.K09.gz43_233465	F	M00056781B:E01	UC2-PrimColon
2077	727737	1839.A11.gz43_233487	F	M00056767B:C01	UC2-PrimColon
2078	477053	1839.F12.gz43_233508	F	M00056775B;H07	UC2-PrimColon
2079	726384	1839.P12.gz43_233518	F	M00056787C:G01	UC2-PrimColon
2080	736686	1839.A13.gz43_233519	F	M00056767B:G08	UC2-PrimColon
2081	483549	1839.L13.gz43_233530	F	M00056782D:F02	UC2-PrimColon
2082	549801	1839,A14.gz43_233535	F	M00056767C:A09	UC2-PrimColon
2083	730484	1839.H14.gz43_233542	F	M00056777D:D04	UC2-PrimColon
2084	647991	1839.J14.gz43_233544	F	M00056780C:H12	UC2-PrimColon
2085	490393	1839,L14.gz43_233546	F	M00056782D:F08	UC2-PrimColon
2086	643968	1839.N14.gz43_233548	F	M00056785C:B09	UC2-PrimColon
2087	641968	1839.B15.gz43_233552	F	M00056770C:A07	UC2-PrimColon
2088	726261	1839.L15.gz43_233562	F	M00056782D:F10	UC2-PrimColon
2089	733891	1839.M15.gz43_233563	F	M00056784B:A01	UC2-PrimColon
2090	730022	1839.O15.gz43_233565	F	M00056786D:A03	UC2-PrimColon
2091	730296	1839.M16.gz43_233579	F	M00056784B:C02	UC2-PrimColon
2092	733149	1839.M17.gz43_233595	F	M00056784B:D06	UC2-PrimColon
2093	728487	1839.A18.gz43_233599	F	M00056768A:C10	UC2-PrimColon
2094	726229	1839.C18.gz43_233601	F	M00056771D:D10	UC2-PrimColon
2095	523868	1839.G18.gz43_233605	F	M00056776D:B02	UC2-PrimColon
2096	577305	1839.L18.gz43_233610	F	M00056783A;C08	UC2-PrimColon
2097	727216	1839.J19.gz43_233624	F	M00056780D:G05	UC2-PrimColon
2098	47461	1839.L19.gz43_233626	F	M00056783B:B01	UC2-PrimColon
2099	550780	1839.N19.gz43_233628	F	M00056785D:C09	UC2-PrimColon
2100	606076	1839.P21.gz43_233662	F	M00056788A;D06	UC2-PrimColon

Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
2101	691229	1839.C22.gz43_233665	F	M00056772A:C08	UC2-PrimColon
2102	727366	1839.D22.gz43 233666	F	M00056773A:G10	UC2-PrimColon
2103	724722	1839.J22.gz43_233672	F	M00056780D:H10	UC2-PrimColon
2104	725784	1839.M22.gz43_233675	F	M00056784C:C06	UC2-PrimColon
2105	732345	1839.B23.gz43_233680	F	M00056771A:F03	UC2-PrimColon
2106	726699	1839.C23.gz43_233681	F	M00056772A:D03	UC2-PrimColon
2107	513156	1839.J23.gz43_233688	F	M00056781A:C05	UC2-PrimColon
2108	631038	1839.E24.gz43_233699	F	M00056774D:B02	UC2-PrimColon
2109	609459	1839.H24.gz43_233702	F	M00056778C;G08	UC2-PrimColon
2110	482788	1839.J24.gz43_233704	F	M00056781A:D02	UC2-PrimColon
2111	730592	1839.N24.gz43_233708	F	M00056785D:G07	UC2-PrimColon
2112	735801	1852.A13.gz43_235742	F	M00056871C:D05	UC2-PrimColon
2113	472704	1852.H13.gz43_235749	F	M00056879D:D01	UC2-PrimColon
2114	649299	1852.I13.gz43_235750	F	M00056881A:A10	UC2-PrimColon
2115	650944	1852.N13.gz43_235755	F	M00056886B:B10	UC2-PrimColon
2116	481641	1852.E14.gz43_235762	F	M00056876C:B02	UC2-PrimColon
2117	488349	1852.F14.gz43_235763	F	M00056877B:H09	UC2-PrimColon
2118	735096	1852.I14.gz43_235766	F	M00056881A:C02	UC2-PrimColon
2119	594434	1852.J14.gz43_235767	F	M00056882B:E12	UC2-PrimColon
2120	550673	1852.L14.gz43_235769	F	M00056884C:H08	UC2-PrimColon
2121	735672	1852.N14.gz43_235771	F	M00056886B:C05	UC2-PrimColon
2122	732712	1852.K15.gz43_235784	F	M00056883D:A07	UC2-PrimColon
2123	472307	1852.L15.gz43_235785	F	M00056884C:H111	UC2-PrimColon
2124	729981	1852.D16.gz43_235793	F	M00056875D:C04	UC2-PrimColon
2125	639950	1852.I16.gz43_235798	F	M00056881A:H02	UC2-PrimColon
2126	724517	1852.L16.gz43_235801	F	M00056884D:C07	UC2-PrimColon
2127	735412	1852.B17.gz43_235807	F	M00056873A:H06	UC2-PrimColon
2128	724223	1852.N17.gz43_235819	F	M00056886B:H02	UC2-PrimColon
2129	727602	1852.D18.gz43_235825	F	M00056875D:E09	UC2-PrimColon
2130	561396	1852.L18.gz43_235833	F	M00056884D:D06	UC2-PrimColon
2131	726408	1852.B19.gz43_235839	F	M00056873B:C09	UC2-PrimColon
2132	730059	1852.C19.gz43_235840	F	M00056874C:D05	UC2-PrimColon
2133	655327	1852.F19.gz43_235843	F	M00056877C:G12	UC2-PrimColon
2134	732872	1852.M19.gz43_235850	F	M00056886A:A09	UC2-PrimColon
2135	473512	1852.N19.gz43_235851	F	M00056886C:D02	UC2-PrimColon
2136	730441	1852.F20.gz43_235859	F	M00056877C:H03	UC2-PrimColon
2137	451184	1852.N20.gz43_235867	F	M00056886C:D11	UC2-PrimColon
2138	733040	1852.D21.gz43_235873	F	M00056875D:H12	UC2-PrimColon
2139	732872	1852.K21.gz43_235880	F	M00056883D:F07	UC2-PrimColon
2140	152	1852.M21.gz43_235882	F	M00056886A:C11	UC2-PrimColon
2141	730301	1852.I22.gz43_235894	F	M00056881B:G04	UC2-PrimColon
2142	471931	1852.J22.gz43_235895	F	M00056882D:A06	UC2-PrimColon

Table 2

	able 2				
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	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
2143	730106	1852.L23.gz43_235913	F	M00056885A:D12	UC2-PrimColon
2144	649030	1852.024.gz43_235932	F	M00056888A:H04	UC2-PrimColon
2145	562345	1861.I04.gz43_235990	F	M00056900C:B12	UC2-PrimColon
2146	472226	1861.P04.gz43_235997	F	M00056910B:F01	UC2-PrimColon
2147	494393	1861.E07.gz43_236034	F	M00056895C:E11	UC2-PrimColon
2148	730393	1861.B08.gz43_236047	F	M00056891C:H08	UC2-PrimColon
2149	676221	1861.G10.gz43_236084	F	M00056898C:B06	UC2-PrimColon
2150	648774	1861.O11.gz43_236108	F	M00056909B:E11	UC2-PrimColon
2151	448368	1861.H12.gz43_236117	F	M00056899D:B06	UC2-PrimColon
2152	489001	1861.K12.gz43_236120	F	M00056902D:H09	UC2-PrimColon
2153	557525	1861.J14.gz43_236151	F	M00056902A:G12	UC2-PrimColon
2154	724773	1861.C16.gz43_236176	F	M00056893B:G12	UC2-PrimColon
2155	730608	1861.I16.gz43_236182	F	M00056901B:C03	UC2-PrimColon
2156	642897	1861.K16.gz43_236184	F	M00056903A:C08	UC2-PrimColon
2157	548275	1861.N16.gz43_236187	F	M00056908B:D02	UC2-PrimColon
2158	728640	1861.C17.gz43_236192	F	M00056893C:A02	UC2-PrimColon
2159	479572	1861.B18.gz43_236207	F	M00056892B:C09	UC2-PrimColon
2160	735834	1861.E18.gz43_236210	F	M00056896A:C01	UC2-PrimColon
2161	733425	1861.K18.gz43_236216	F	M00056903A:F03	UC2-PrimColon
2162	732653	1861.P19.gz43_236237	F	M00056911B:A01	UC2-PrimColon
2163	729060	1861.G20.gz43_236244	F	M00056898D:H09	UC2-PrimColon
2164	656667	1861,H21.gz43_236261	F	M00056900B:D01	UC2-PrimColon
2165	732676	1861.G22.gz43_236276	F	M00056899A:A11	UC2-PrimColon
2166	730296	1861.H24.gz43_236309	F	M00056900B:F07	UC2-PrimColon
2167	453001	1861.K24.gz43_236312	F	M00056903B:D09	UC2-PrimColon
2168	725451	1861,N24.gz43_236315	F	M00056908D:A11	UC2-PrimColon
2169	730851	1863.C01.gz43_236704	F	M00056936C:E04	UC2-PrimColon
2170	727596	1863.H01.gz43_236709	F	M00056941D:D07	UC2-PrimColon
2171	736293	1863.L01.gz43_236713	F	M00056946A:F07	UC2-PrimColon
2172	733910	1863.N01.gz43_236715	F	M00056948C:F03	UC2-PrimColon
2173	77737	1863.C02.gz43_236720	F	M00056936C:F11	UC2-PrimColon
2174	638983	1863.L02.gz43_236729	F	M00056946A:G06	UC2-PrimColon
2175	736001	1863.M02.gz43_236730	F	M00056947C:B04	UC2-PrimColon
2176	731196	1863.P02.gz43_236733	F	M00056950D:F12	UC2-PrimColon
2177	446820	1863.A03.gz43_236734	F	M00056934A:E07	UC2-PrimColon
2178	734151	1863.E03.gz43_236738	F	M00056939A:C01	UC2-PrimColon
2179	456183	1863.H03.gz43_236741	F	M00056941D:G05	UC2-PrimColon
2180	734629	1863.I03.gz43_236742	F	M00056943A:B04	UC2-PrimColon
2181	733623	1863.J03.gz43_236743	F	M00056944A:D09	UC2-PrimColon
2182	733856	1863.N03.gz43_236747	F	M00056948D:A11	UC2-PrimColon
2183	727480	1863.P03.gz43_236749	F	M00056950D:H09	UC2-PrimColon
2184	448233	1863.J04.gz43_236759	F	M00056944B:C02	UC2-PrimColon

Table 2

	able 2				
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ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
2185	729691	1863.I05.gz43 236774	F	M00056943A:F01	UC2-PrimColon
2186	500239	1863.N05.gz43 236779	F	M00056949A:A04	UC2-PrimColon
2187	646314	1863.K06.gz43 236792	F	M00056945A:F02	UC2-PrimColon
2188	492893	1863.A08.gz43 236814	F	M00056934C:C04	UC2-PrimColon
2189	451624	1863.D08.gz43 236817	F	M00056938A:H09	UC2-PrimColon
2190	552416	1863.L08.gz43 236825	F	M00056946C:B08	UC2-PrimColon
2191	555478	1863.A10.gz43 236846	F	M00056934C:G09	UC2-PrimColon
2192	616541	1863.G10.gz43 236852	F	M00056941B:B02	UC2-PrimColon
2193	410487	1863.O10.gz43 236860	F	M00056950A:H03	UC2-PrimColon
2194	726825	1863.B11.gz43 236863	F	M00056935C:G08	UC2-PrimColon
2195	733144	1863.D12.gz43 236881	F	M00056938B:C09	UC2-PrimColon
2196	642528	1863.E12.gz43 236882	F	M00056939B:E05	UC2-PrimColon
2197	735994	1863.J12.gz43 236887	F	M00056944C:B03	UC2-PrimColon
2198	724401	1863.K12.gz43 236888	F	M00056945C:A11	UC2-PrimColon
2199	736129	1863.M12.gz43 236890	F	M00056948A:D09	UC2-PrimColon
2200	448046	1863.P13.gz43_236909	F	M00056951B:F09	UC2-PrimColon
2201	731216	1863.D15.gz43_236929	F	M00056938B:H08	UC2-PrimColon
2202	449061	1863.M15.gz43 236938	F	M00056948B:B03	UC2-PrimColon
2203	730815	1863.F17.gz43 236963	F	M00056940C:E05	UC2-PrimColon
2204	732756	1863.H17.gz43 236965	F	M00056942C:F11	UC2-PrimColon
2205	468015	1863.I17.gz43 236966	F	M00056943C:A01	UC2-PrimColon
2206	733081	1863.J17.gz43_236967	F	M00056944D:A06	UC2-PrimColon
2207	555484	1863.L17.gz43 236969	F	M00056946D:G09	UC2-PrimColon
2208	731355	1863.A18.gz43_236974	F	M00056935A:C02	UC2-PrimColon
2209	730375	1863.G18.gz43_236980	F	M00056941B:G08	UC2-PrimColon
2210	467803	1863.M18.gz43_236986	F	M00056948B:G05	UC2-PrimColon
2211	735306	1863.D20.gz43_237009	F	M00056938C:F10	UC2-PrimColon
2212	729316	1863.I20.gz43_237014	F	M00056943D:B02	UC2-PrimColon
2213	551441	1863.L20.gz43_237017	F	M00056947A:C05	UC2-PrimColon
2214	735484	1863.A21.gz43_237022	F	M00056935A:F04	UC2-PrimColon
2215	171511	1863.E21.gz43_237026	F	M00056939D:B02	UC2-PrimColon
2216	728076	1863.H21.gz43_237029	F	M00056942D:D07	UC2-PrimColon
2217	511351	1863.H22.gz43_237045	F	M00056942D:D11	UC2-PrimColon
2218	731125	1863.I22.gz43_237046	F	M00056943D:H08	UC2-PrimColon
2219	524706	1863.B24.gz43_237071	F	M00056936C:C06	UC2-PrimColon
2220	485880	1863.C24.gz43_237072	F	M00056937C:H08	UC2-PrimColon
2221	736197	1864.I03.gz43_237126	F	M00056961C:C07	UC2-PrimColon
2222	731089	1864.M04.gz43_237146	F	M00056968C:C06	UC2-PrimColon
2223	642781	1864.G12.gz43_237268	F	M00056959C:B10	UC2-PrimColon
2224	731947	1864.M14.gz43_237306	F	M00056969A:B07	UC2-PrimColon
2225	448712	1864.M16.gz43_237338	F	M00056969A:C07	UC2-PrimColon
2226	736289	1864.I17.gz43_237350	F	M00056962D:F09	UC2-PrimColon

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SEQ ID NO	CLUSTER	SEQ NAME	ORIEN T	CLONE ID	LIBRARY
2227	735123	1873.E02.gz43 237490	F	M00056978D:B07	UC2-PrimColon
2228	724411	1873.H02.gz43 237493	F	M00056982D:B12	UC2-PrimColon
2229	662617	1873.P02.gz43 237501	F	M00056993D:C05	UC2-PrimColon
2230	639934	1873.M03.gz43 237514	F	M00056989C:H09	UC2-PrimColon
2231	733563	1873.C04.gz43_237520	F	M00056977A:C02	UC2-PrimColon
2232	203793	1873.K04.gz43 237528	F	M00056987C:D08	UC2-PrimColon
2233	703978	1873.L04.gz43 237529	F	M00056988C:D05	UC2-PrimColon
2234	732114	1873.P05.gz43 237549	F	M00056993D:F05	UC2-PrimColon
2235	732213	1873.J06.gz43_237559	F	M00056986A:E09	UC2-PrimColon
2236	725825	1873.N06.gz43 237563	F	M00056990D:C11	UC2-PrimColon
2237	734936	1873.D08.gz43 237585	F	M00056978A:A03	UC2-PrimColon
2238	731607	1873.E08.gz43_237586	F	M00056979B:D03	UC2-PrimColon
2239	731302	1873.K10.gz43_237624	F	M00056987D:A09	UC2-PrimColon
2240	732113	1873.L10.gz43_237625	F	M00056988D:F05	UC2-PrimColon
2241	728013	1873.M10.gz43_237626	F	M00056989D:F07	UC2-PrimColon
2242	730089	1873.D11.gz43_237633	F	M00056978A:H04	UC2-PrimColon
2243	648034	1873.E12.gz43_237650	F	M00056979C:D11	UC2-PrimColon
2244	729623	1873.I12.gz43_237654	F	M00056985A:D06	UC2-PrimColon
2245	731317	1873.M12.gz43_237658	F	M00056989D:H11	UC2-PrimColon
2246	728249	1873.A13.gz43_237662	F	M00056975A:H11	UC2-PrimColon
2247	613029	1873.C13.gz43_237664	F	M00056977B:B06	UC2-PrimColon
2248	731542	1873.O13.gz43_237676	F	M00056993A:B08	UC2-PrimColon
2249	732535	1873.G14.gz43_237684	F	M00056981D:H02	UC2-PrimColon
2250	735440	1873.A16.gz43_237710	F	M00056975B:E04	UC2-PrimColon
2251	641658	1873.D16.gz43_237713	F	M00056978B:G02	UC2-PrimColon
2252	406931	1873.N18.gz43_237755	F	M00056991C:H11	UC2-PrimColon
2253	736401	1873.P18.gz43_237757	_ F	M00056994B:H05	UC2-PrimColon
2254	734561	1873.B20.gz43_237775	F	M00056976C:F05	UC2-PrimColon
2255	727151	1873.F20.gz43_237779	F	M00056980D:E07	UC2-PrimColon
2256	731453	1873.I20.gz43_237782	F	M00056985C:C06	UC2-PrimColon
2257	642649	1873.K20.gz43_237784	F	M00056988A:F06	UC2-PrimColon
2258	642288	1873.M20.gz43_237786	F	M00056990B:H07	UC2-PrimColon
2259	572426	1873.P20.gz43_237789	F	M00056994C:B04	UC2-PrimColon
2260	731358	1873,K22.gz43_237816	F	M00056988B:A06	UC2-PrimColon
2261	471883	1873.N22.gz43_237819	F	M00056992A;E01	UC2-PrimColon
2262	733778	1873.D24.gz43_237841	F	M00056978D:A01	UC2-PrimColon
2263	731592	1873.J24.gz43_237847	F	M00056987A:C02	UC2-PrimColon
2264	456236	1874.D01.gz43_237857	F	M00056998C:B10	UC2-PrimColon
2265	732792	1874.N01.gz43_237867	F	M00057009C:B02	UC2-PrimColon
2266	733932	1874.E03.gz43_237890	F	M00057000A:A05	UC2-PrimColon
2267	724810	1874.P07.gz43_237965	F	M00057011C:H03	UC2-PrimColon
2268	731317	1874.B08.gz43_237967	F	M00056996D:A02	UC2-PrimColon

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SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
2269	726408	1874.D08.gz43 237969	F	M00056998D:H08	UC2-PrimColon
2270	734708	1874.D12.gz43 238033	F	M00056999A:G12	UC2-PrimColon
2271	631251	1874.A15.gz43 238078	F	M00056995C:H06	UC2-PrimColon
2272	726696	1874.D16.gz43 238097	F	M00056999B:D07	UC2-PrimColon
2273	448202	1874.F16.gz43 238099	F	M00057001D:F02	UC2-PrimColon
2274	734629	1874.H16.gz43_238101	F	M00057003D:F02	UC2-PrimColon
2275	561993	1874.K16.gz43 238104	F	M00057007B;G02	UC2-PrimColon
2276	407723	1874.J17.gz43 238119	F	M00057006A:G10	UC2-PrimColon
2277	736413	1874.K17.gz43_238120	F	M00057007C:A06	UC2-PrimColon
2278	556637	1874.P17.gz43 238125	F	M00057012A:D12	UC2-PrimColon
2279	731686	1874.A19.gz43 238142	F	M00056995D:C11	UC2-PrimColon
2280	559053	1874.B20.gz43_238159	F	M00056997B:C11	UC2-PrimColon
2281	733625	1875.J01.gz43 238247	F	M00057024B:F07	UC2-PrimColon
2282	461486	1875.O01.gz43 238252	F	M00057030C;B03	UC2-PrimColon
2283	556019	1875.B02.gz43 238255	F	M00057014B:A02	UC2-PrimColon
2284	732119	1875.F02.gz43 238259	F	M00057018C:F02	UC2-PrimColon
2285	411128	1875.K02.gz43_238264	F	M00057025C:A08	UC2-PrimColon
2286	732441	1875.A03.gz43_238270	F	M00057012D:G03	UC2-PrimColon
2287	478448	1875.B03.gz43_238271	F	M00057014B:B01	UC2-PrimColon
2288	733723	1875.D03.gz43_238273	F	M00057016B:A09	UC2-PrimColon
2289	726173	1875.I03.gz43_238278	F	M00057023A:H09	UC2-PrimColon
2290	451351	1875.B04.gz43_238287	F	M00057014B:B06	UC2-PrimColon
2291	570573	1875.E04.gz43_238290	F	M00057017A:F11	UC2-PrimColon
2292	655312	1875.K04.gz43_238296	F	M00057025C:D11	UC2-PrimColon
2293	732047	1875.C05.gz43_238304	F	M00057015A:E02	UC2-PrimColon
2294	734091	1875.F05.gz43_238307	F	M00057018D:B10	UC2-PrimColon
2295	538582	1875.E06.gz43_238322	F	M00057017A:G04	UC2-PrimColon
2296	487183	1875.C07.gz43_238336	F	M00057015A:G06	UC2-PrimColon
2297	483919	1875.G07.gz43_238340	F	M00057020D:A05	UC2-PrimColon
2298	651049	1875.L07.gz43_238345	F	M00057027B:B11	UC2-PrimColon
2299	734392	1875.D08.gz43_238353	F	M00057016B:E06	UC2-PrimColon
2300	557896	1875.F08.gz43_238355	F	M00057018D:E05	UC2-PrimColon
2301	89082	1875.L08.gz43_238361	F	M00057027B:E04	UC2-PrimColon
2302	512863	1875.P08.gz43_238365	F	M00057032A;C01	UC2-PrimColon
2303	474009	1875.D09.gz43_238369	F	M00057016B:H08	UC2-PrimColon
2304	729731	1875.H09.gz43_238373	F	M00057022B;A04	UC2-PrimColon
2305	481581	1875.A10.gz43_238382	F	M00057013B:H07	UC2-PrimColon
2306	732550	1875.C10.gz43_238384	F	M00057015A:H12	UC2-PrimColon
2307	735396	1875.J10.gz43_238391	F	M00057024D:D12	UC2-PrimColon
2308	642693	1875.P10.gz43_238397	F	M00057032A:F12	UC2-PrimColon
2309	733970	1875.D11.gz43_238401	F	M00057016C:B05	UC2-PrimColon
2310	639629	1875.H11.gz43_238405	F	M00057022B:F03	UC2-PrimColon

Table 2

	able 2				
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ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
2311	731577	1875.L11.gz43 238409	F	M00057027B:F06	UC2-PrimColon
2312	733354	1875.B12.gz43_238415	F	M00057014C:E01	UC2-PrimColon
2313	626791	1875.F12.gz43_238419	F	M00057019A:G04	UC2-PrimColon
2314	559699	1875.H12.gz43_238421	F	M00057022B:H04	UC2-PrimColon
2315	726208	1875.A13.gz43_238430	F	M00057013C:E09	UC2-PrimColon
2316	730828	1875.B13.gz43_238431	F	M00057014C:E03	UC2-PrimColon
2317	390968	1875.I13.gz43_238438	F	M00057023D:D05	UC2-PrimColon
2318	471522	1875.O13.gz43_238444	F	M00057031A:H02	UC2-PrimColon
2319	731838	1875.D14.gz43_238449	F	M00057016C:E04	UC2-PrimColon
2320	726476	1875.H14.gz43_238453	F	M00057022C:D12	UC2-PrimColon
2321	466920	1875.I14.gz43_238454	F	M00057023D:D08	UC2-PrimColon
2322	732712	1875.N14.gz43_238459	F	M00057029D:A06	UC2-PrimColon
2323	561338	1875.O14.gz43_238460	F	M00057031B:A01	UC2-PrimColon
2324	733756	1875.A15.gz43 238462	F	M00057013C:H01	UC2-PrimColon
2325	484023	1875.C15.gz43_238464	F	M00057015B:E01	UC2-PrimColon
2326	557262	1875.D15.gz43_238465	F	M00057016C:E11	UC2-PrimColon
2327	559096	1875.E15.gz43_238466	F	M00057017C:E03	UC2-PrimColon
2328	455148	1875.F15.gz43_238467	F	M00057019C:C08	UC2-PrimColon
2329	616985	1875.I15.gz43_238470	F	M00057023D:E11	UC2-PrimColon
2330	733479	1875.N15.gz43_238475	F	M00057029D:F01	UC2-PrimColon
2331	558134	1875.F16.gz43_238483	F	M00057019C:E03	UC2-PrimColon
2332	703298	1875.D17.gz43_238497	F	M00057016D:B07	UC2-PrimColon
2333	477797	1875.G18.gz43_238516	F	M00057021B:B07	UC2-PrimColon
2334	432159	1875.L18.gz43_238521	F	M00057027D:A12	UC2-PrimColon
2335	736385	1875.P18.gz43_238525	F	M00057032D:A04	UC2-PrimColon
2336	619265	1875.B19.gz43_238527	F	M00057014D:E05	UC2-PrimColon
2337	4255	1875.F19.gz43_238531	F	M00057019D:C02	UC2-PrimColon
2338	554080	1875.M19.gz43_238538	F	M00057028D:D09	UC2-PrimColon
2339	734074	1875.C21.gz43_238560	F	M00057016A:B04	UC2-PrimColon
2340	734177	1875.E21.gz43_238562	F	M00057018A:C05	UC2-PrimColon
2341	727761	1875.I21.gz43_238566	F	M00057024A:D08	UC2-PrimColon
2342	733209	1875.L21.gz43_238569	F	M00057027D:D07	UC2-PrimColon
2343	512377	1875.A22.gz43_238574	F	M00057014A;B11	UC2-PrimColon
2344	733664	1875.B22.gz43_238575	F	M00057014D:G10	UC2-PrimColon
2345	640904	1875.D22.gz43_238577	F	M00057016D:H03	UC2-PrimColon
2346	425203	1875.N22.gz43_238587	F	M00057030B:B03	UC2-PrimColon
2347	557164	1875.B23.gz43_238591	F	M00057014D:H01	UC2-PrimColon
2348	494450	1875.C23.gz43_238592	F	M00057016A:G01	UC2-PrimColon
2349	639427	1875.F23.gz43_238595	F	M00057020A;F09	UC2-PrimColon
2350	554693	1875.H23.gz43_238597	F	M00057022D:E10	UC2-PrimColon
2351	732598	1875.K23.gz43_238600	F	M00057026D:A05	UC2-PrimColon
2352	495241	1875.M23.gz43_238602	F	M00057029A:C08	UC2-PrimColon

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	able 2				
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ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
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2354	726044	1875.I24.gz43 238614	F	M00057024B;A11	UC2-PrimColon
2355	728791	1875.L24.gz43 238617	F	M00057027D;G03	UC2-PrimColon
2356	562769	1875.N24.gz43 238619	F	M00057030C:A05	UC2-PrimColon
2357	479208	1875.P24.gz43 238621	F	M00057033A:D08	UC2-PrimColon
2358	4244	1876.C03.gz43 238656	F	M00057035C:E10	UC2-PrimColon
2359	732159	1876.K03.gz43 238664	F	M00057046A:F02	UC2-PrimColon
2360	473026	1876.E04.gz43 238674	F	M00057038A:H07	UC2-PrimColon
2361	736385	1876.B05.gz43 238687	F	M00057034C:G12	UC2-PrimColon
2362	734466	1876.J05.gz43 238695	F	M00057045A:C04	UC2-PrimColon
2363	668731	1876.K05.gz43 238696	F	M00057046A:G05	UC2-PrimColon
2364	736881	1876.H06.gz43 238709	F	M00057042D:E06	UC2-PrimColon
2365	466920	1876.O06.gz43 238716	F	M00057051B:E09	UC2-PrimColon
2366	728353	1876.E07.gz43 238722	F	M00057038C:B06	UC2-PrimColon
2367	732246	1876.N07.gz43 238731	F	M00057049D:F12	UC2-PrimColon
2368	468296	1876.A08.gz43 238734	F	M00057033C:B03	UC2-PrimColon
2369	454819	1876.E09.gz43 238754	F	M00057038C:G08	UC2-PrimColon
2370	481293	1876.M10.gz43 238778	F	M00057048C:E04	UC2-PrimColon
2371	735815	1876.E11.gz43 238786	F	M00057038D:A12	UC2-PrimColon
2372	732315	1876.H12.gz43 238805	F	M00057043A:G07	UC2-PrimColon
2373	558573	1876.M12.gz43 238810	F	M00057048C:H11	UC2-PrimColon
2374	492627	1876,G13.gz43 238820	F	M00057041D:C08	UC2-PrimColon
2375	732042	1876.K14.gz43 238840	F	M00057046C:E05	UC2-PrimColon
2376	737087	1876.N14.gz43_238843	F	M00057050B:F06	UC2-PrimColon
2377	456469	1876.A16.gz43_238862	F	M00057033D:F10	UC2-PrimColon
2378	735871	1876.I16.gz43_238870	F	M00057044C:B05	UC2-PrimColon
2379	736855	1876.F18.gz43_238899	F	M00057040D:H04	UC2-PrimColon
2380	732562	1876.M20.gz43_238938	F.	M00057048D:H10	UC2-PrimColon
2381	727321	1876.G22.gz43_238964	·F	M00057042B:A10	UC2-PrimColon
2382	638857	1876.H22.gz43_238965	F	M00057043C:H11	UC2-PrimColon
2383	735972	1876.A23.gz43_238974	F	M00057034B:B01	UC2-PrimColon
2384	735054	1876.L23.gz43_238985	F	M00057047D:H04	UC2-PrimColon
2385	554854	1885.M01.gz43_239018	F	M00057068D:F04	UC2-PrimColon
2386	731262	1885.O01.gz43_239020	F	M00057071A:A10	UC2-PrimColon
2387	729981	1885.C02.gz43_239024	F	M00057055C:F01	UC2-PrimColon
2388	735292	1885.F02.gz43_239027	F	M00057059D:H09	UC2-PrimColon
2389	726699	1885.L02.gz43_239033	F	M00057067D:H06	UC2-PrimColon
2390	624133	1885.O02.gz43_239036	F	M00057071A:B04	UC2-PrimColon
2391	645262	1885,A03.gz43_239038	F	M00057053A:A02	UC2-PrimColon
2392	558549	1885,K03.gz43_239048	F	M00057066D:B03	UC2-PrimColon
2393	736093	1885.L03.gz43_239049	F	M00057068A:C10	UC2-PrimColon
2394	733573	1885.G04.gz43_239060	· F	M00057061B:H02	UC2-PrimColon

Table 2

	able 2				
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2396	556385	1885.N04.gz43_239067	F	M00057070A:B07	UC2-PrimColon
2397	559656	1885.L05.gz43_239081	F	M00057068A:F05	UC2-PrimColon
2398	446616	1885.A06.gz43 239086	F	M00057053A:D11	UC2-PrimColon
2399	728445	1885.C06.gz43_239088	F	M00057055C:H07	UC2-PrimColon
2400	732223	1885.D06.gz43_239089	F	M00057056D:F11	UC2-PrimColon
2401	595506	1885.G06.gz43_239092	F	M00057061C:D04	UC2-PrimColon
2402	627515	1885.K06.gz43_239096	F	M00057066D:E01	UC2-PrimColon
2403	730528	1885.L06.gz43_239097	F	M00057068A:F07	UC2-PrimColon
2404	451233	1885.E07.gz43_239106	F	M00057058D:G08	UC2-PrimColon
2405	728884	1885.H07.gz43_239109	F	M00057063A:C08	UC2-PrimColon
2406	541793	1885.I07.gz43_239110	F	M00057064C:F11	UC2-PrimColon
2407	736415	1885.L08.gz43_239129	F	M00057068A:G05	UC2-PrimColon
2408	552641	1885.F09.gz43_239139	F	M00057060B:D07	UC2-PrimColon
2409	736156	1885.G09.gz43_239140	F	M00057061D:D03	UC2-PrimColon
2410	560581	1885.M09.gz43_239146	F	M00057069A:F09	UC2-PrimColon
2411	654723	1885.D10.gz43_239153	F	M00057057A:G01	UC2-PrimColon
2412	731449	1885.E10.gz43_239154	F	M00057059A:B04	UC2-PrimColon
2413	452623	1885.G10.gz43_239156	F	M00057061D:F05	UC2-PrimColon
2414	732072	1885.M10.gz43_239162	F	M00057069A:H08	UC2-PrimColon
2415	452224	1885.P10.gz43_239165	F	M00057072C:A03	UC2-PrimColon
2416	550637	1885.K11.gz43_239176	F	M00057067B;C11	UC2-PrimColon
2417	453508	1885.M11.gz43 239178	F	M00057069B:A08	UC2-PrimColon
2418	732872	1885.A12.gz43_239182	F	M00057053C:B10	UC2-PrimColon
2419	735283	1885.P12.gz43_239197	F	M00057072C:A09	UC2-PrimColon
2420	422590	1885.D13.gz43_239201	F	M00057057B:E06	UC2-PrimColon
2421	422242	1885.J13.gz43_239207	F	M00057066A:A09	UC2-PrimColon
2422	471982	1885.D14.gz43_239217	F	M00057057B:E07	UC2-PrimColon
2423	736285	1885.H14.gz43_239221	F	M00057063B:F06	UC2-PrimColon
2424	734787	1885.P14.gz43_239229	F	M00057072C:C02	UC2-PrimColon
2425	733623	1885.A15.gz43_239230	F	M00057053C:G04	UC2-PrimColon
2426	736318	1885.D15.gz43_239233	F	M00057057B:G03	UC2-PrimColon
2427	677769	1885.A16.gz43_239246	F	M00057053D:D01	UC2-PrimColon
2428	726380	1885.B16.gz43_239247	_ F	M00057055B:D07	UC2-PrimColon
2429	134501	1885.C16.gz43_239248	F	M00057056B:C06	UC2-PrimColon
2430	731844	1885.F16.gz43_239251	F	M00057060C:D05	UC2-PrimColon
2431	730472	1885.B17.gz43_239263	F	M00057055B:E10	UC2-PrimColon
2432	455884	1885.C17.gz43_239264	F	M00057056B:D05	UC2-PrimColon
2433	669159	1885.H17.gz43_239269	F	M00057063C:C04	UC2-PrimColon
2434	736146	1885.K17.gz43_239272	F	M00057067C:D04	UC2-PrimColon
2435	729502	1885.A18.gz43_239278	F	M00057054A:B12	UC2-PrimColon
2436	731830	1885.C18.gz43_239280	F	M00057056B:D11	UC2-PrimColon

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Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
2437	706245	1885.N18.gz43_239291	F	M00057070C:H10	UC2-PrimColon
2438	631526	1885.P18.gz43_239293	F	M00057072C:H01	UC2-PrimColon
2439	736113	1885.A19.gz43_239294	F	M00057054A:D01	UC2-PrimColon
2440	567005	1885.B19.gz43_239295	F	M00057055B:F05	UC2-PrimColon
2441	730128	1885.K19.gz43_239304	F	M00057067C:H09	UC2-PrimColon
2442	620462	1885.E20.gz43_239314	F	M00057059C:H02	UC2-PrimColon
2443	450551	1885,F20.gz43_239315	F	M00057060D:C09	UC2-PrimColon
2444	724773	1885.L20,gz43_239321	F	M00057068D:B03	UC2-PrimColon
2445	735028	1885.N20.gz43_239323	F	M00057070D:B08	UC2-PrimColon
2446	476947	1885.E21.gz43_239330	F	M00057059D:A09	UC2-PrimColon
2447	420958	1885.I21.gz43_239334	F	M00057065C:B07	UC2-PrimColon
2448	447758	1885.O21.gz43_239340	F	M00057071D:E01	UC2-PrimColon
2449	447075	1885.G22.gz43_239348	F	M00057062B:H04	UC2-PrimColon
2450	455248	1885.L22.gz43_239353	F	M00057068D:C09	UC2-PrimColon
2451	732351	1885.O22.gz43_239356	F	M00057071D:G01	UC2-PrimColon
2452	620159	1885.P22.gz43_239357	F	M00057073A:B12	UC2-PrimColon
2453	472704	1885.K23.gz43_239368	F	M00057067D:F03	UC2-PrimColon
2454	649349	1885.N23.gz43_239371	F	M00057070D:G03	UC2-PrimColon
2455	735216	1885.F24.gz43_239379	F	M00057061A:F09	UC2-PrimColon
2456	572273	1885.J24.gz43_239383	F	M00057066C:B02	UC2-PrimColon
2457	736210	1885.L24.gz43_239385	F	M00057068D:E05	UC2-PrimColon
2458	731358	1886.M02.gz43_239418	F	M00057088B:E02	UC2-PrimColon
2459	639629	1886.D03.gz43_239425	F	M00057077B:B06	UC2-PrimColon
2460	733910	1886.L03.gz43_239433	F	M00057087B:A07	UC2-PrimColon
2461	473588	1886,N04,gz43_239451	F	M00057090A:B02	UC2-PrimColon
2462	727407	1886.N05.gz43_239467	F	M00057090A:C03	UC2-PrimColon
2463	368965	1886.H07.gz43_239493	F	M00057082C:G03	UC2-PrimColon
2464	734059	1886.L13.gz43_239593	F	M00057087D:B04	UC2-PrimColon
2465	419465	1886.P15.gz43_239629	F	M00057092C:A05	UC2-PrimColon
2466	732487	1886.A18.gz43_239662	F	M00057073D:H05	UC2-PrimColon
2467	538582	1886.M20.gz43_239706	F	M00057089B;D01	UC2-PrimColon
2468	732131	1886.O22.gz43_239740	F	M00057091D:F11	UC2-PrimColon
2469	556212	1886.E23.gz43_239746	F	M00057079A:F05	UC2-PrimColon
2470	731966	1886.M24.gz43_239770	F	M00057089D:E03	UC2-PrimColon
2471	732873	1887.C02.gz43_239809	F	M00057095B:G03	UC2-PrimColon
2472	732550	1887.G02.gz43_239813	F	M00057099D:D11	UC2-PrimColon
2473	731477	1887.D04.gz43_239842	F	M00057096C:B08	UC2-PrimColon
2474	485441	1887.I04.gz43_239847	F	M00057102A:F12	UC2-PrimColon
2475	533689	1887,B05.gz43_239856	F	M00057094B:D07	UC2-PrimColon
2476	735515	1887.H05.gz43_239862	F	M00057100D:E09	UC2-PrimColon
2477	678846	1887,M06.gz43_239883	F	M00057106A:H04	UC2-PrimColon
2478	638908	1887.P06.gz43_239886	F	M00057110C:A04	UC2-PrimColon

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SEQ ID NO CLUSTER SEQ NAME T CLONE ID LIBRARY	T	able 2				
2479 516799 1887.L07.gz43 239898 F M00057105C:C06 UC2-PrimColon		CLUSTER	SEQ NAME		CLONE ID	LIBRARY
2480 553898 1887.A08_pz43 239903 F M00057093C.A02 UC2-PrimColon 2481 644354 1887.F09_pz43 239924 F M00057093A.C04 UC2-PrimColon 2482 731697 1887.F10_pz43 239929 F M00057093A.P02 UC2-PrimColon 2484 732885 1887.F10_pz43 239940 F M00057109A.P11 UC2-PrimColon 2485 736634 1887.F10_pz43 239951 F M00057095C.E01 UC2-PrimColon 2486 457842 1887.D11_pz43 239951 F M00057095C.E01 UC2-PrimColon 2487 654475 1887.F11_pz43 239954 F M00057095C.E01 UC2-PrimColon 2488 454772 1887.F11_pz43 239954 F M00057095D.H05 UC2-PrimColon 2488 454772 1887.F11_pz43 239957 F M00057095D.H05 UC2-PrimColon 2489 734344 1887.L12_pz43 239978 F M00057103E.E12 UC2-PrimColon 2490 693869 1887.N12_pz43 239980 F M00057103E.E12 UC2-PrimColon 2491 733348 1887.O12_pz43 249010 F M00057105D.H09 UC2-PrimColon 2491 733348 1887.O12_pz43 240010 F M00057105D.H09 UC2-PrimColon 2493 401368 1887.F15_pz43 240020 F M00057105D.H09 UC2-PrimColon 2494 452276 1887.H17_pz43 24005 F M00057105D.H09 UC2-PrimColon 2494 452276 1887.H17_pz43 24005 F M00057105D.H0 UC2-PrimColon 2496 735240 1887.N12_pz43 240060 F M00057105D.H0 UC2-PrimColon 2496 735254 1887.D19_pz43 240069 F M00057105D.H0 UC2-PrimColon 2496 735254 1887.D19_pz43 240069 F M00057105C.F0 UC2-PrimColon 2499 474387 1887.C0_pz43 240090 F M00057105C.F0 UC2-PrimColon 2499 47387 1887.C0_pz43 240090 F M00057105C.F0 UC2-PrimColon 2496 735212 1887.D19_pz43 240090 F M00057105C.F0 UC2-PrimColon 2500 733913 1887.E12_pz43 240110 F M00057105A.F11 UC2-PrimColon 2500 735220 1887.P10_pz43 240110 F M00057105A.F11 UC2-PrimColon 2500 73525 1888.D10_pz43 240115 F M00057105A.F11 UC2-PrimColon 2500 73525 1888.D10_pz43 240115 F M00057105A.F11 UC2-PrimColon 2500 73525 1888.D10_pz43	2479	516799	1887 I 07 gr43 239898	F		
2481 644354 1887.F09.gz43 239924 F M00057099A.C04 UC2-PrimColon 2482 731697 1887.K09.gz43 239940 F M0005709A.H12 UC2-PrimColon 2483 645139 1887.F10.gz43 239940 F M0005709A.H12 UC2-PrimColon 2484 732885 1887.O10.gz43 239940 F M0005709A.H12 UC2-PrimColon 2485 736634 1887.A11.gz43 239954 F M00057095.H12 UC2-PrimColon 2486 457842 1887.D11.gz43 239954 F M0005709D.H05 UC2-PrimColon 2487 654475 1887.F11.gz43 239956 F M0005709D.H05 UC2-PrimColon 2488 343772 1887.H1.gz43 239956 F M0005709D.H05 UC2-PrimColon 2489 734344 1887.H1.gz43 239978 F M00057102B.E12 UC2-PrimColon 2490 693869 1887.N12.gz43 239980 F M00057103C.602 UC2-PrimColon 2491 733348 1887.O12.gz43 239981 F M00057103D.F10 UC2-PrimColon 2492 7333209 1887.H1.gz43 239981 F M00057105D.F10 UC2-PrimColon 2494 452276 1887.H1.gz43 240020 F M00057105D.F10 UC2-PrimColon 2494 452276 1887.H1.gz43 240020 F M00057105D.F10 UC2-PrimColon 2494 452276 1887.H1.gz43 240069 F M00057105D.F10 UC2-PrimColon 2496 735584 1887.G18.gz43 240069 F M00057105D.F10 UC2-PrimColon 2496 735584 1887.G18.gz43 240069 F M00057105D.F10 UC2-PrimColon 2498 733212 1887.O19.gz49 240097 F M00057100C.F02 UC2-PrimColon 2498 733212 1887.O19.gz43 240098 F M00057100C.F02 UC2-PrimColon 2498 733212 1887.O19.gz43 240098 F M00057100C.F02 UC2-PrimColon 2498 733212 1887.D19.gz43 240098 F M00057106C.F02 UC2-PrimColon 2500 735131 1888.D01.gz43 24010 F M00057103A.F11 UC2-PrimColon 2500 735131 1888.D01.gz43 240112 F M00057103A.F11 UC2-PrimColon 2500 735131 1888.D01.gz43 240175 F M00057112A.C12 UC2-PrimColon 2500 736548 1888.D01.gz43 240175 F M00057112A.C12 UC2-PrimColon 2500 736548 1888.D01.gz43 240175 F M00057112D.F06 UC2-PrimColon 2510 602673 1888.D01.gz43						
2482 731697 1887.K09.gz43 239929 F M00057104D-A02 UC2-PrimColon 2483 645139 1887.F10.gz43 239940 F M00057099A-H12 UC2-PrimColon 2484 732885 1887.O10.gz43 239951 F M00057099A-H12 UC2-PrimColon 2485 736634 1887.A11.gz43 239951 F M0005709B-M0 UC2-PrimColon 2486 457842 1887.D11.gz43 239956 F M0005709B-M0 UC2-PrimColon 2487 654475 1887.F11.gz43 239956 F M0005709B-M0 UC2-PrimColon 2488 543772 1887.112.gz43 239956 F M0005709B-M0 UC2-PrimColon 2498 734344 1887.L12.gz43 23995 F M00057102B-E12 UC2-PrimColon 2490 693869 1887.N12.gz43 239980 F M00057103B-A12 UC2-PrimColon 2490 693869 1887.N12.gz43 239981 F M00057103B-A12 UC2-PrimColon 2492 733209 1887.L12.gz43 239981 F M00057103B-A12 UC2-PrimColon 2493 401368 1887.F15.gz43 240020 F M00057103D-C10 UC2-PrimColon 2493 401368 1887.F15.gz43 240020 F M00057103D-C10 UC2-PrimColon 2495 736220 1887.P17.gz43 240055 F M00057102D-C11 UC2-PrimColon 2495 736220 1887.P17.gz43 240062 F M00057102D-C11 UC2-PrimColon 2496 735384 1887.G18.gz43 240069 F M00057102D-C11 UC2-PrimColon 2496 735384 1887.G18.gz43 240088 F M00057109C-D10 UC2-PrimColon 2498 732312 1887.D19.gz43 240098 F M00057109C-D10 UC2-PrimColon 2498 732312 1887.D19.gz43 240098 F M00057109C-D10 UC2-PrimColon 2498 732312 1887.D19.gz43 240098 F M00057109C-D10 UC2-PrimColon 2500 732922 1887.D20.gz43 240098 F M00057109C-D10 UC2-PrimColon 2500 735313 1887.B21.gz43 240115 F M00057109C-D10 UC2-PrimColon 2500 73553 1888.D01.gz43 240115 F M00057109C-D10 UC2-PrimColon 2504 725960 1887.D2.gz43 240115 F M00057103A-G11 UC2-PrimColon 2504 725960 1887.D2.gz43 240115 F M00057103A-G11 UC2-PrimColon 2506 735256 1888.D01.gz43 240260 F M00057112A-G08 UC2-PrimColon 2506 736539 1888.D01.gz4						
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2495 736220 1887,P17,g243_240062 F M00057110D:E12 UC2-PrimColon 2496 735584 1887,G18.gz/43_240068 F M00057104A-H06 UC2-PrimColon 2497 448325 1887,U19.gz/43_240088 F M00057104A-H06 UC2-PrimColon 2498 732312 1887,O19.gz/43_240093 F M00057109C:D10 UC2-PrimColon 2499 477387 1887,C20.gz/43_240093 F M00057096E:O5 UC2-PrimColon 2500 729222 1887,D20.gz/43_240098 F M0005709FE:D2 UC2-PrimColon 2501 733972 1887,F20.gz/43_240100 F M0005709FE:D2 UC2-PrimColon 2502 735113 1887,E21.gz/43_240112 F M00057095E:D2 UC2-PrimColon 2503 659020 1887,E21.gz/43_240119 F M00057095A:F11 UC2-PrimColon 2504 729560 1887,O24.gz/43_240173 F M00057110A:A03 UC2-PrimColon 2505 415825 1888,A01.gz/43_240173 F M00057110A:A03 UC2-PrimColon 2506 733225 1888,D10.gz/43_240175 F M00057112A:C12 UC2-PrimColon 2506 733225 1888,D10.gz/43_240178 F M00057112A:C12 UC2-PrimColon 2508 568632 1888,L03.gz/43_240218 F M00057112A:G08 UC2-PrimColon 2509 736539 1888,D02.gz/43_240218 F M00057112A:G08 UC2-PrimColon 2510 602673 1888,D05.gz/43_240242 F M00057112A:G08 UC2-PrimColon 2511 553264 1888,F06.gz/43_240260 F M00057112A:G08 UC2-PrimColon 2512 555555 1888,G06.gz/43_240261 F M00057112D:F06 UC2-PrimColon 2514 451764 1888,O06.gz/43_240260 F M00057112D:F01 UC2-PrimColon 2515 732896 1888,D07.gz/43_240279 F M00057112D:H01 UC2-PrimColon 2516 733365 1888,D07.gz/43_240279 F M00057112D:D06 UC2-PrimColon 2516 733365 1888,D07.gz/43_240279 F M00057112D:D06 UC2-PrimColon 2517 736548 1888,D09.gz/43_240279 F M00057112D:D06 UC2-PrimColon 2518 53338 1888,D09.gz/43_240301 F M00057112D:D06 UC2-PrimColon 2519 105056 1888,B07.gz/43_240301 F M00057112D:D06 UC2-PrimColon 2519 105056 1888,B07.gz/43_240301 F M00057112D:D06 UC2-PrimColon 2519 105056 1888,B07.gz/43_240301 F M000						
2496						
2497 448325 1887,J19,g243 240088 F M00057104A;H06 UC2-PrimColon 2498 732312 1887,O19,g243 240097 F M000571096;CD10 UC2-PrimColon 2499 477387 1887,C20,g243 240098 F M000570978;EO3 UC2-PrimColon 2500 729222 1887,D20,g243 240098 F M000570978;EO3 UC2-PrimColon 2502 733113 1887,B21,g243 240112 F M000570995;AF11 UC2-PrimColon 2503 650920 1887,D21,g243 240112 F M00057095A;F11 UC2-PrimColon 2504 729560 1887,D21,g243 240113 F M00057103A;F11 UC2-PrimColon 2504 729560 1887,D21,g243 240113 F M00057103A;F11 UC2-PrimColon 2504 729560 1887,D21,g243 240173 F M00057110A;A03 UC2-PrimColon 2506 733225 1888,D01,g243 240178 F M00057115D;C10 UC2-PrimColon 2506 733225 1888,D01,g243 240194 F M00057115D;C10 UC2-PrimColon 2509 736539 1888,D4,g243 24018 F M00057112A;C12 UC2-PrimColon 2509 736539 1888,D4,g243 240218 F M0005712A;D06 UC2-PrimColon 2510 6062673 1888,D05,g243 240242 F M00057115D;F00 UC2-PrimColon 2511 553264 1888,F06,g243 240260 F M0005712A;D10 UC2-PrimColon 2512 555655 1888,G06,g243 240266 F M0005712D;D60 UC2-PrimColon 2513 735634 1888,H03,7g243 240266 F M0005712D;D10 UC2-PrimColon 2514 451764 1888,D07,g243 240266 F M0005712D;D10 UC2-PrimColon 2515 732896 1888,D07,g243 240269 F M0005712D;D10 UC2-PrimColon 2515 733365 1888,D07,g243 240279 F M0005712D;D10 UC2-PrimColon 2516 53338 1888,D07,g243 240297 F M0005712D;D10 UC2-PrimColon 2516 53338 1888,D07,g243 240299 F M0005712D;D10 UC2-PrimColon 2516 53338 1888,D07,g243						
2498 732312 1887.O19.gz43 240093 F M00057109C:D10 UC2-PrimColon						
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2500 729222 1887.D20_gz43_240098 F M00057097B:E02 UC2-PrimColon 2501 733972 1887.P20_gz43_240100 F M00057099C:A06 UC2-PrimColon 2502 735113 1887.D21_gz43_240112 F M00057099C:A06 UC2-PrimColon 2503 650920 1887.D21_gz43_240115 F M00057103A:F11 UC2-PrimColon 2504 729560 1887.D21_gz43_240115 F M00057103A:F11 UC2-PrimColon 2505 415825 1888.A01_gz43_240175 F M00057110A:A03 UC2-PrimColon 2506 733225 1888.D01_gz43_240176 F M00057112A:C12 UC2-PrimColon 2506 733225 1888.D01_gz43_240178 F M00057113D:C10 UC2-PrimColon 2507 732113 1888.D02_gz43_240194 F M00057115D:D06 UC2-PrimColon 2508 568632 1888.D32_gz43_240218 F M00057115D:D06 UC2-PrimColon 2509 736539 1888.D32_gz43_240232 F M00057115D:F06 UC2-PrimColon 2510 602673 1888.D05_gz43_240224 F M00057115D:F06 UC2-PrimColon 2511 553264 1888.F06_gz43_240266 F M00057112A:D01 UC2-PrimColon 2512 555555 1888.G06_gz43_240261 F M0005712D:D10 UC2-PrimColon 2514 451764 1888.O06_gz43_240265 F M0005712D:F03 UC2-PrimColon 2515 736534 1888.K06_gz43_240265 F M0005712D:F03 UC2-PrimColon 2516 733365 1888.I07_gz43_240274 F M0005712D:F03 UC2-PrimColon 2516 733365 1888.I07_gz43_240279 F M0005712D:A01 UC2-PrimColon 2517 736548 1888.K08_gz43_240310 F M0005712D:A01 UC2-PrimColon 2517 736548 1888.K08_gz43_240310 F M0005712D:A01 UC2-PrimColon 2518 553338 1888.K08_gz43_240310 F M0005712D:B110 UC2-PrimColon 2518 553338 1888.K08_gz43_240310 F M0005712D:B110 UC2-PrimColon 2518 553338 1888.K08_gz43_240310 F M0005712D:B110 UC2-PrimColon 2518 553338 1888.K08_gz43_240329 F M0005712D:C0 UC2-PrimColon 2518 553338 1888.K08_gz43_240310 F M0005712D:C0 UC2-PrimColon 2518 553338 1888.K08_gz43_240310 F M0005712D:C0 UC2-PrimColon 2518 553538 1888.K08_gz43_240329 F M0005712D:C0 UC2-PrimColon 2518						
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2502 735113 1887.B21.gz43 240112 F M00057095A:F11 UC2-PrimColon 2503 650920 1887.L21.gz43 240119 F M00057103A:F11 UC2-PrimColon 2504 729560 1887.U21.gz43 240113 F M00057103A:F11 UC2-PrimColon 2505 415825 1888.A01.gz43 240175 F M00057112A:C12 UC2-PrimColon 2506 733225 1888.D01.gz43 240175 F M00057112A:C12 UC2-PrimColon 2507 732113 1888.D02.gz43 240194 F M00057115D:D06 UC2-PrimColon 2509 736539 1888.L03.gz43 240218 F M00057112D:D06 UC2-PrimColon 2510 602673 1888.D05.gz43 240242 F M00057113D:P06 UC2-PrimColon 2510 602673 1888.D05.gz43 240240 F M00057113D:P06 UC2-PrimColon 2511 553264 1888.F06.gz43 240240 F M00057113D:P06 UC2-PrimColon 2512 555655 1888.G06.gz43 240260 F M00057112D:D06 UC2-PrimColon 2513 736534 1888.G06.gz43 240269 F M0005712D:D:F03 UC2-PrimColon 2514 451764 1888.D06.gz43 240269 F M0005712D:D:F03 UC2-PrimColon 2515 73286 1888.D07.gz43 240269 F M0005712D:D:F03 UC2-PrimColon 2516 733365 1888.D07.gz43 240279 F M0005712D:D:D1 UC2-PrimColon 2516 733365 1888.D07.gz43 240279 F M0005712D:D:D1 UC2-PrimColon 2516 733365 1888.D07.gz43 240279 F M0005712D:D:D1 UC2-PrimColon 2516 53338 1888.H09.gz43 240301 F M0005712D:D:D1 UC2-PrimColon 2516 53338 1888.H09.gz43 240301 F M0005712D:D:D1 UC2-PrimColon 2517 736548 1888.D07.gz43 240301 F M0005712D:D:D1 UC2-PrimColon 2518 53338 1888.H09.gz43 240301 F M0005712D:D:D1 UC2-PrimColon 2519 105056 1888.K10.gz43 240309 F M0005712D:D:D0 UC2-PrimColon 2519 105056 1						
2503 650920 1887.121.g243 240119 F M00057103A:F11 UC2-PrimColon	2502	735113				
2504 729560 1887.024.gz43 240173 F M00057110A:A03 UC2-PrimColon	2503	650920				
2505 415825 1888.A01.gz43 240175 F M00057112A:C12 UC2-PrimColon	2504	729560		F		
2506 733225 1888.D01.gz43 240178 F M00057115D:C10 UC2-PrimColon	2505					
2507 732113 1888.D02.gz43 240194 F M00057115D-D06 UC2-PrimColon	2506			F		
2509 736539 1888,104 gz43 240232 F M00057124A:G08 UC2-PrimColon 2510 602673 1888,D05,gz43 240242 F M00057115D:F06 UC2-PrimColon 2511 553264 1888,F06,gz43 240260 F M00057118B:E10 UC2-PrimColon 2512 555655 1888,G06,gz43 240261 F M00057120A:D01 UC2-PrimColon 2513 736534 1888,K06,gz43 240265 F M00057120A:D11 UC2-PrimColon 2514 451764 1888,D06,gz43 240269 F M0005712D:F03 UC2-PrimColon 2515 7332896 1888,D07,gz43 240274 F M0005712D:F03 UC2-PrimColon 2516 73365 1888,I07,gz43 240279 F M0005712D:A01 UC2-PrimColon 2517 736548 1888,O08,gz43 240301 F M0005712D:A01 UC2-PrimColon 2518 553338 1888,H09,gz43 240310 F M0005712D:H01 UC2-PrimColon 2518 553338 1888,H09,gz43 240310 F M0005712D:H01 UC2-PrimColon	2507	732113	1888.D02,gz43 240194	F	M00057115D:D06	
2509 736539 1888,104,gz43 240232 F M00057124A:G08 UC2-PrimColon 2510 602673 1888,D05,gz43 240242 F M00057115D:F06 UC2-PrimColon 2511 553264 1888,F06,gz43 240260 F M00057118B:E10 UC2-PrimColon 2512 555655 1888,G06,gz43 240261 F M00057120A:D01 UC2-PrimColon 2513 736534 1888,K06,gz43 240265 F M00057120A:D01 UC2-PrimColon 2514 451764 1888,D07,gz43 240274 F M00057112D:F03 UC2-PrimColon 2516 733365 1888,I07,gz43 240279 F M0005712D:A01 UC2-PrimColon 2516 73365 1888,I09,gz43 240301 F M0005712D:A01 UC2-PrimColon 2518 53338 1888,H09,gz43 240310 F M0005712D:H01 UC2-PrimColon 2518 53338 1888,H09,gz43 240310 F M0005712D:H01 UC2-PrimColon 2518	2508	568632	1888.L03.gz43 240218	F	M00057126D:A04	UC2-PrimColon
2511 553264 1888 F06 gz/43 240260 F M00057118E:E10 UC2-PrimColon 2512 555655 1888 G06 gz/43 240261 F M00057120A:D01 UC2-PrimColon 2513 736548 1888 K06 gz/3 240265 F M00057120C:B11 UC2-PrimColon 2514 451764 1888 O06 gz/43 240269 F M00057120D:F03 UC2-PrimColon 2515 73286 1888 D07 gz/43 240279 F M0005712D:A01 UC2-PrimColon 2516 733365 1888 U07 gz/43 240279 F M0005712D:A01 UC2-PrimColon 2517 736548 1888 O08 gz/43 240301 F M0005712D:A01 UC2-PrimColon 2518 553338 1888 H09 gz/43 240310 F M0005712D:H01 UC2-PrimColon 2519 105056 1888 K10 gz/43 240329 F M0005712D:E02 UC2-PrimColon	2509	736539	1888.J04.gz43 240232	F	M00057124A:G08	UC2-PrimColon
2512 555655 1888.G06_gz43 240261 F M00057120A:D01 UC2-PrimColon 2513 736534 1888.K06_gz43 240265 F M00057125C:B11 UC2-PrimColon 2514 451764 1888.006_gz43 240269 F M00057129D:F03 UC2-PrimColon 2515 732896 1888.D07_gz43 240274 F M00057116A:B06 UC2-PrimColon 2516 733365 1888.107_gz43 240279 F M0005712D:A01 UC2-PrimColon 2517 736548 1888.008_gz43 240310 F M0005712B:H01 UC2-PrimColon 2518 553338 1888.H09_gz43 240310 F M0005712B:H10 UC2-PrimColon 2519 105056 1888.K10_gz43 240310 F M0005712B:CE02 UC2-PrimColon	2510	602673	1888.D05.gz43 240242	F	M00057115D:F06	UC2-PrimColon
2513 736534 1888.K06.gz/43 240265 F M00057125C:B11 UC2-PrimColon	2511	553264		F	M00057118B;E10	UC2-PrimColon
2514 451764 1888.006.gz43 240269 F M00057129D:F03 UC2-PrimColon	2512	555655	1888.G06.gz43 240261	F	M00057120A;D01	UC2-PrimColon
2515 732896 1888.D07.gz43 240274 F M00057116A:B06 UC2-PrimColon 2516 733365 1888.I07.gz43 240279 F M00057122D:A01 UC2-PrimColon 2517 736548 1888.008.gz43 240310 F M00057130A:A02 UC2-PrimColon 2518 553338 1888.H09.gz43 240310 F M0005712B:H10 UC2-PrimColon 2519 105056 1888.K10.gz43 240329 F M0005712C:E02 UC2-PrimColon	2513	736534	1888,K06,gz43 240265	F	M00057125C:B11	UC2-PrimColon
2515 732896 1888.D07.gz43_240274 F M00057116A:B06 UC2-PrimColon 2516 733365 1888.I07.gz43_240279 F M00057122D:A01 UC2-PrimColon 2517 736548 1888.008.gz43_240301 F M0005712D:A01 UC2-PrimColon 2518 553338 1888.H09.gz43_240310 F M0005712B:H10 UC2-PrimColon 2519 105056 1888.K10.gz43_240329 F M00057125C:E02 UC2-PrimColon	2514	451764	1888.O06.gz43 240269	F	M00057129D:F03	UC2-PrimColon
2517 736548 1888,008,gz43 240301 F M00057130A:A02 UC2-PrimColon 2518 553338 1888,H09,gz43 240310 F M00057121B:H10 UC2-PrimColon 2519 105056 1888,K10,gz43 440329 F M00057125C:E02 UC2-PrimColon	2515	732896		F	M00057116A:B06	UC2-PrimColon
2518 553338 1888.H09.gz43_240310 F M00057121B:H10 UC2-PrimColon 2519 105056 1888.K10.gz43_240329 F M00057125C:E02 UC2-PrimColon	2516	733365	1888.I07.gz43 240279	F	M00057122D:A01	UC2-PrimColon
2518 553338 1888.H09.gz43_240310 F M00057121B:H10 UC2-PrimColon 2519 105056 1888.K10.gz43_240329 F M00057125C:E02 UC2-PrimColon	2517			F		
	2518	553338		F		
	2519	105056	1888.K10.gz43 240329	F	M00057125C:E02	UC2-PrimColon
2020 200030 1000.010.g245_240333 1 MIUUU3/13UA:AII UC2-PIIIICOIOII	2520	288535	1888.O10.gz43_240333	F	M00057130A;A11	UC2-PrimColon

PCT/US01/25840

Table 2

1	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
2521	453733	1888.G12.gz43_240357	F	M00057120C:B09	UC2-PrimColon
2522	552783	1888.H12.gz43_240358	F	M00057121C:E08	UC2-PrimColon
2523	732937	1888.I12.gz43_240359	F	M00057123A:F09	UC2-PrimColon
2524	542490	1888.K12.gz43_240361	F	M00057125D:B02	UC2-PrimColon
2525	553087	1888.L12.gz43_240362	F	M00057127A:E10	UC2-PrimColon
2526	609459	1888.G14.gz43_240389	F	M00057120C:F08	UC2-PrimColon
2527	736810	1888.O14.gz43_240397	F	M00057130A:D10	UC2-PrimColon
2528	735959	1888.L15.gz43_240410	F	M00057127A:H10	UC2-PrimColon
2529	560868	1888.H17.gz43_240438	F	M00057122A:A07	UC2-PrimColon
2530	726176	1888.P17.gz43_240446	F	M00057131D:D02	UC2-PrimColon
2531	724952	1888,A18.gz43_240447	F	M00057112D:G08	UC2-PrimColon
2532	1030	1888.H18.gz43_240454	F	M00057122A:C04	UC2-PrimColon
2533	454961	1888.L18.gz43_240458	F	M00057127B:D10	UC2-PrimColon
2534	643517	1888.C19.gz43_240465	F	M00057115C:B11	UC2-PrimColon
2535	733006	1888.H19.gz43_240470	F	M00057122A:C11	UC2-PrimColon
2536	733171	1888.A20.gz43_240479	F	M00057113A:A08	UC2-PrimColon
2537	735018	1888.B20.gz43_240480	F	M00057114C:E07	UC2-PrimColon
2538	728479	1888.B21.gz43_240496	F	M00057114C:F08	UC2-PrimColon
2539	726575	1888.J21.gz43_240504	F	M00057125A:A07	UC2-PrimColon
2540	734606	1888.I22.gz43_240519	F	M00057123D:F07	UC2-PrimColon
2541	732476	1888.D23.gz43_240530	F	M00057116C:H09	UC2-PrimColon
2542	656263	1888.H23.gz43_240534	F	M00057122B:F04	UC2-PrimColon
2543	662617	1888.K23.gz43_240537	F	M00057126C:C05	UC2-PrimColon
2544	474108	1888.D24.gz43_240546	F	M00057116D:B10	UC2-PrimColon
2545	455581	1888.J24.gz43_240552	F	M00057125A:F07	UC2-PrimColon
2546	447692	1888.L24.gz43_240554	F	M00057127C:F03	UC2-PrimColon
2547	735514	1888.M24.gz43_240555	F	M00057128C:H02	UC2-PrimColon
2548	440284	1888.N24.gz43_240556	F	M00057129D:D07	UC2-PrimColon
2549	639644	1897.A02.gz43_240575	F	M00057132B:H05	UC2-PrimColon
2550	568031	1897.G02.gz43_240581	F	M00057139A:G08	UC2-PrimColon
2551	555960	1897.L02.gz43_240586	F	M00057144D:D03	UC2-PrimColon
2552	462779	1897.M02.gz43_240587	F	M00057145D:E01	UC2-PrimColon
2553	472068	1897.C03.gz43_240593	F	M00057134B:D06	UC2-PrimColon
2554	728413	1897.N03.gz43_240604	F	M00057147A:B07	UC2-PrimColon
2555	519378	1897.P03.gz43_240606	F	M00057149B;B07	UC2-PrimColon
2556	733951	1897.A04.gz43_240607	F	M00057132C:C04	UC2-PrimColon
2557	448212	1897.P04.gz43_240622	F	M00057149B:B08	UC2-PrimColon
2558	536225	1897.B05.gz43_240624	F	M00057133C:B02	UC2-PrimColon
2559	736288	1897.F05.gz43_240628	F	M00057138A:F02	UC2-PrimColon
2560	225960	1897.E06.gz43_240643	F	M00057137A:H12	UC2-PrimColon
2561	655327	1897.H08.gz43_240678	F	M00057140B:H01	UC2-PrimColon
2562	735729	1897.N08.gz43_240684	F	M00057147A:H07	UC2-PrimColon

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Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
2563	642273	1897.F09.gz43 240692	F	M00057138B:B02	UC2-PrimColon
2564	549285	1897.009.gz43 240701	F	M00057148C:B02	UC2-PrimColon
2565	735756	1897.P11.gz43 240734	F	M00057149C:H01	UC2-PrimColon
2566	640662	1897.E12.gz43 240739	F	M00057137B:C07	UC2-PrimColon
2567	543772	1897.F12.gz43 240740	F	M00057138B:H02	UC2-PrimColon
2568	735801	1897.M12.gz43_240747	F	M00057146A;H05	UC2-PrimColon
2569	457846	1897.N12.gz43 240748	F	M00057147C:B01	UC2-PrimColon
2570	736318	1897.O12.gz43_240749	F	M00057148C:C09	UC2-PrimColon
2571	658271	1897.K13.gz43_240761	F	M00057144B:D04	UC2-PrimColon
2572	735283	1897.M13.gz43_240763	F	M00057146B:C06	UC2-PrimColon
2573	447550	1897.A15.gz43_240783	F	M00057133A:B07	UC2-PrimColon
2574	500758	1897.H15.gz43_240790	F	M00057140D:B03	UC2-PrimColon
2575	567005	1897.H17.gz43_240822	F	M00057140D:F02	UC2-PrimColon
2576	523171	1897.K17.gz43 240825	F	M00057144B:H10	UC2-PrimColon
2577	397515	1897.M17.gz43_240827	F	M00057146B:H12	UC2-PrimColon
2578	736318	1897.F18.gz43_240836	F	M00057138D:F03	UC2-PrimColon
2579	449035	1897.I18.gz43_240839	F	M00057142A:H07	UC2-PrimColon
2580	486076	1897.K18.gz43_240841	F	M00057144C:A02	UC2-PrimColon
2581	729843	1897.O18.gz43_240845	F	M00057149A:A04	UC2-PrimColon
2582	726440	1897.F19.gz43_240852	F	M00057138D:F10	UC2-PrimColon
2583	731317	1897.G19.gz43_240853	F	M00057139D:G07	UC2-PrimColon
2584	559004	1897.I22.gz43_240903	F	M00057142B:F06	UC2-PrimColon
2585	448202	1897.L22.gz43_240906	F	M00057145C:H03	UC2-PrimColon
2586	718314	1897.N22.gz43_240908	F	M00057147D:H09	UC2-PrimColon
2587	735989	1897.C23.gz43_240913	F	M00057135A:B02	UC2-PrimColon
2588	489001	1897.F23.gz43_240916	F	M00057139A:B10	UC2-PrimColon
2589	462779	1897.H23.gz43_240918	F	M00057141A:G06	UC2-PrimColon
2590	596809	1898.C02.gz43_242213	F	M00057153B:G07	UC2-PrimColon
2591	447003	1898.A03.gz43_242227	F	M00057150D:C04	UC2-PrimColon
2592	730661	1898.D04.gz43 242246	F	M00057155A:E11	UC2-PrimColon
2593	461316	1898.H04.gz43_242250	F	M00057160D:A11	UC2-PrimColon
2594	450724	1898.A05.gz43_242259	F	M00057150D:F08	UC2-PrimColon
2595	480307	1898.B05.gz43_242260	F	M00057152B:H02	UC2-PrimColon
2596	727608	1898.F05.gz43_242264	F	M00057157D:H08	UC2-PrimColon
2597	649852	1898.P05.gz43_242274	F	M00057169A:F05	UC2-PrimColon
2598	650297	1898.A06.gz43_242275	F	M00057151A:B04	UC2-PrimColon
2599	491933	1898.E06.gz43_242279	F	M00057156B:D10	UC2-PrimColon
2600	733552	1898.P06.gz43_242290	F	M00057169A:F06	UC2-PrimColon
2601	730341	1898,D07.gz43_242294	F	M00057155A:G11	UC2-PrimColon
2602	640563	1898.D08.gz43_242310	F	M00057155A:H07	UC2-PrimColon
2603	640603	1898.B10.gz43_242340	F	M00057152C:C10	UC2-PrimColon
2604	552972	1898.E10.gz43_242343	F	M00057156C:E08	UC2-PrimColon

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Table 2

	able 2				
SEQ ID NO	CI LIGHED	gno.v.v	ORIEN		
	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
2605	736030	1898.F10.gz43_242344	F	M00057158B:C02	UC2-PrimColon
2606	736228	1898.G11.gz43_242361	F	M00057159C:E11	UC2-PrimColon
2607	648820	1898.M11.gz43_242367	F	M00057165D:H05	UC2-PrimColon
2608	735993	1898.D12.gz43_242374	_ F	M00057155C:B07	UC2-PrimColon
2609	402516	1898.H12.gz43_242378	F	M00057161B:D02	UC2-PrimColon
2610	583625	1898.O12.gz43_242385	F	M00057168B:D02	UC2-PrimColon
2611	726448	1898.N13.gz43_242400	F	M00057167A:D08	UC2-PrimColon
2612	735756	1898.H14.gz43_242410	F	M00057161B:E07	UC2-PrimColon
2613	551654	1898.C15.gz43_242421	F	M00057153D:H01	UC2-PrimColon
2614	558477	1898.D15.gz43_242422	F	M00057155C:G04	UC2-PrimColon
2615	728353	1898.115.gz43_242427	F	M00057162B:E10	UC2-PrimColon
2616	736634	1898.M15.gz43_242431	F	M00057166B:B07	UC2-PrimColon
2617	737006	1898.P15.gz43_242434	F	M00057169C:H10	UC2-PrimColon
2618	736288	1898.J16.gz43_242444	_ F	M00057163C:D06	UC2-PrimColon
2619	734794	1898.P16.gz43_242450	F	M00057169C:H12	UC2-PrimColon
2620	734928	1898.E17.gz43_242455	F	M00057156D:F02	UC2-PrimColon
2621	727976	1898.F17.gz43_242456	F	M00057158C:C10	UC2-PrimColon
2622	736790	1898.F19.gz43_242488	_ F	M00057158C:G05	UC2-PrimColon
2623	735633	1898.I21.gz43_242523	F	M00057162C:C01	UC2-PrimColon
2624	734484	1898.B22.gz43_242532	F	M00057153B:A04	UC2-PrimColon
2625	480142	1898.D22.gz43_242534	F	M00057155D:E12	UC2-PrimColon
2626	454073	1898.G22.gz43_242537	F	M00057160B:G08	UC2-PrimColon
2627	599714	1898.L22.gz43_242542	F	M00057165C:E08	UC2-PrimColon
2628	737042	1898.M22.gz43_242543	F	M00057166B:G06	UC2-PrimColon
2629	405016	1898.N22.gz43_242544	F	M00057167C:A05	UC2-PrimColon
2630	723915	1898.L23.gz43_242558	_ F	M00057165C:F05	UC2-PrimColon
2631	737006	1898.O23.gz43_242561	F	M00057168C:D03	UC2-PrimColon
2632	734808	1898.B24.gz43_242564	F	M00057153B:D02	UC2-PrimColon
2633	471887	1898.P24.gz43_242578	F	M00057170A:D11	UC2-PrimColon
2634	733972	1899.A01.gz43_242579	F	M00057170A:D06	UC2-PrimColon
2635	554221	1899.J01.gz43_242588	F	M00057181C:D10	UC2-PrimColon
2636	735426	1899.N01.gz43_242592	F	M00057192A:D12	UC2-PrimColon
2637	454812	1899.P01.gz43_242594	F	M00057194C:A03	UC2-PrimColon
2638	562453	1899.D02.gz43_242598	F	M00057173D:B12	UC2-PrimColon
2639	554683	1899.M02.gz43_242607	F	M00057189D:G08	UC2-PrimColon
2640	474346	1899.N02.gz43_242608	F	M00057192A:F01	UC2-PrimColon
2641	729039	1899.D03.gz43_242614	F	M00057173D:C07	UC2-PrimColon
2642	560183	1899.J03.gz43_242620	F	M00057181C:G07	UC2-PrimColon
2643	422590	1899.D04.gz43_242630	F	M00057173D:E04	UC2-PrimColon
2644	642936	1899.K04.gz43_242637	F	M00057182C:C03	UC2-PrimColon
2645	730106	1899.N04.gz43_242640	F	M00057192A:H05	UC2-PrimColon
2646	449548	1899.H05.gz43_242650	F	M00057179C:B05	UC2-PrimColon

Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
2647	736861	1899.P05.gz43_242658	F	M00057194C:F02	UC2-PrimColon
2648	730220	1899.A06.gz43_242659	F	M00057170A:H04	UC2-PrimColon
2649	736309	1899.I06.gz43_242667	F	M00057180C:F09	UC2-PrimColon
2650	373615	1899.K06.gz43_242669	F	M00057182C:C11	UC2-PrimColon
2651	640635	1899.006.gz43_242673	F	M00057193C:F11	UC2-PrimColon
2652	735764	1899.F07.gz43_242680	F	M00057176A:H10	UC2-PrimColon
2653	735801	1899.J07.gz43_242684	F	M00057181D:C09	UC2-PrimColon
2654	477098	1899.K07.gz43 242685	F	M00057182D:A08	UC2-PrimColon
2655	734031	1899.M07.gz43_242687	F	M00057191A:H09	UC2-PrimColon
2656	419711	1899.N07.gz43_242688	F	M00057192B:D10	UC2-PrimColon
2657	390968	1899.P07.gz43_242690	F	M00057194C:F10	UC2-PrimColon
2658	732625	1899.E08.gz43_242695	F	M00057175B:G12	UC2-PrimColon
2659	462986	1899.F08.gz43_242696	F	M00057176A:H12	UC2-PrimColon
2660	482512	1899.H08.gz43 242698	F	M00057179C:G10	UC2-PrimColon
2661	473238	1899.C09.gz43 242709	F	M00057172D:F08	UC2-PrimColon
2662	735789	1899.I09.gz43 242715	F	M00057180D:C10	UC2-PrimColon
2663	736595	1899.K09.gz43 242717	F	M00057182D:B11	UC2-PrimColon
2664	735514	1899.M09.gz43 242719	F	M00057191B;C06	UC2-PrimColon
2665	630516	1899.O09.gz43 242721	F	M00057193D:F06	UC2-PrimColon
2666	730899	1899.N10.gz43 242736	F	M00057192C:B11	UC2-PrimColon
2667	552055	1899.P10.gz43 242738	F	M00057194D:C12	UC2-PrimColon
2668	528404	1899.C11.gz43 242741	F	M00057173A:C07	UC2-PrimColon
2669	734209	1899.D11.gz43 242742	F	M00057174B:C06	UC2-PrimColon
2670	452243	1899.J11.gz43 242748	F	M00057181D:H07	UC2-PrimColon
2671	734061	1899.O11.gz43 242753	F	M00057194A:B07	UC2-PrimColon
2672	625988	1899.G12.gz43 242761	F	M00057177C:D07	UC2-PrimColon
2673	561632	1899.D13.gz43 242774	F	M00057174B:C11	UC2-PrimColon
2674	731370	1899.E13.gz43 242775	F	M00057175C:D02	UC2-PrimColon
2675	480142	1899.G13.gz43 242777	F	M00057177C:G11	UC2-PrimColon
2676	736049	1899.O13.gz43 242785	F	M00057194A:C06	UC2-PrimColon
2677	725691	1899.A14.gz43 242787	F	M00057170C:C01	UC2-PrimColon
2678	727523	1899.M14.gz43 242799	F	M00057191C:D02	UC2-PrimColon
2679	421794	1899.D15.gz43 242806	F	M00057174B:F11	UC2-PrimColon
2680	732429	1899.G15.gz43 242809	F	M00057177C:H04	UC2-PrimColon
2681	530883	1899.H15.gz43_242810	F	M00057180A:H11	UC2-PrimColon
2682	729173	1899.P15.gz43 242818	F	M00057196A:E03	UC2-PrimColon
2683	734744	1899.D16.gz43 242822	F	M00057174B:G03	UC2-PrimColon
2684	735605	1899.F16.gz43 242824	F	M00057176B:F10	UC2-PrimColon
2685	726786	1899.N16.gz43 242832	F	M00057192D:G02	UC2-PrimColon
2686	725089	1899.O16.gz43 242833	F	M00057194A:F01	UC2-PrimColon
2687	448770	1899.D17.gz43 242838	F	M00057174B:G12	UC2-PrimColon
2688	532904	1899.E17.gz43 242839	F	M00057175D:A12	UC2-PrimColon

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Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	Т	CLONE ID	LIBRARY
2689	560568	1899.K17.gz43 242845	F	M00057186A:E12	UC2-PrimColon
2690	736402	1899.N17.gz43_242848	F	M00057192D:G04	UC2-PrimColon
2691	516729	1899.P17.gz43_242850	F	M00057196B:H10	UC2-PrimColon
2692	718314	1899.H18.gz43_242858	F	M00057180B:C06	UC2-PrimColon
2693	737116	1899.J18.gz43_242860	F	M00057182A:H07	UC2-PrimColon
2694	723985	1899.B19.gz43_242868	F	M00057172A:B02	UC2-PrimColon
2695	573733	1899.F19.gz43_242872	F	M00057176C:H08	UC2-PrimColon
2696	475872	1899.H19.gz43_242874	F	M00057180B:D01	UC2-PrimColon
2697	625988	1899.F20.gz43_242888	F	M00057176D:A03	UC2-PrimColon
2698	736354	1899.L20.gz43_242894	F	M00057189C:G11	UC2-PrimColon
2699	535866	1899.C21.gz43_242901	F	M00057173C:C07	UC2-PrimColon
2700	648221	1899.D21.gz43_242902	F	M00057174C:G07	UC2-PrimColon
2701	736280	1899.H21.gz43_242906	F	M00057180B:F05	UC2-PrimColon
2702	156329	1899.N21.gz43_242912	F	M00057193B:C11	UC2-PrimColon
2703	721768	1899.021.gz43_242913	F	M00057194B:E05	UC2-PrimColon
2704	639507	1899.B22.gz43_242916	F	M00057172A:H06	UC2-PrimColon
2705	451383	1899.E22.gz43_242919	F	M00057175D:E12	UC2-PrimColon
2706	349977	1899.J22.gz43_242924	F	M00057182B:D09	UC2-PrimColon
2707	734928	1899.M22.gz43_242927	F	M00057191D:G10	UC2-PrimColon
2708	480623	1899.A23.gz43_242931	F	M00057170D:B08	UC2-PrimColon
2709	674526	1899.H23.gz43_242938	F	M00057180B:G06	UC2-PrimColon
2710	730805	1899.M23.gz43_242943	F	M00057192A:B04	UC2-PrimColon
2711	640350	1900.G01.gz43_242969	F	M00057206A:C06	UC2-PrimColon
2712	726449	1900.K01.gz43_242973	F	M00057210A:C12	UC2-PrimColon
2713	552457	1900.E02.gz43_242983	F	M00057203B:F08	UC2-PrimColon
2714	730308	1900.C04.gz43_243013	F	M00057200D:E03	UC2-PrimColon
2715	734993	1900.E04.gz43_243015	F	M00057203C:A09	UC2-PrimColon
2716	562320	1900.H04.gz43_243018	F	M00057207B:F06	UC2-PrimColon
2717	556656	1900.O04.gz43_243025	F	M00057216D:D05	UC2-PrimColon
2718	446752	1900.D06.gz43_243046	F	M00057202A:D05	UC2-PrimColon
2719	736738	1900.L06.gz43_243054	F	M00057211C:C06	UC2-PrimColon
2720	639395	1900.007.gz43_243073	F	M00057216D:F10	UC2-PrimColon
2721	516729	1900.B08.gz43_243076	F	M00057199B:B06	UC2-PrimColon
2722	557262	1900.E08.gz43_243079	F	M00057203D:A09	UC2-PrimColon
2723	731531	1900.I08.gz43_243083	F	M00057208B:H08	UC2-PrimColon
2724	736778	1900.K08.gz43_243085	F	M00057210C:D09	UC2-PrimColon
2725	724638	1900.010.gz43_243121	F	M00057217A:A12	UC2-PrimColon
2726	483061	1900.B12.gz43_243140	F	M00057199C:H08	UC2-PrimColon
2727	725905	1900.C12.gz43_243141	F	M00057201A:H03	UC2-PrimColon
2728	648757	1900,M12.gz43_243151	F	M00057213C:D06	UC2-PrimColon
2729	735797	1900.A14.gz43_243171	F	M00057197D;H10	UC2-PrimColon
2730	732771	1900.I14.gz43_243179	F	M00057208C:E12	UC2-PrimColon

Table 2

	able 2				
SEO			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CI ONE ID	******
				CLONE ID	LIBRARY
2731	736276	1900.E15.gz43_243191	F	M00057204A:F11	UC2-PrimColon
2732	449078	1900.O19.gz43_243265	F	M00057217B:F12	UC2-PrimColon
2733	161489	1900.A20.gz43_243267	F	M00057198B:C02	UC2-PrimColon
2734	734226	1900.F20.gz43_243272	F	M00057205D:G06	UC2-PrimColon
2735	736500	1900.G20.gz43_243273	F	M00057207A;A07	UC2-PrimColon
2736	733272	1900,K20,gz43_243277	F	M00057211A:D01	UC2-PrimColon
2737	532904	1900.B22.gz43_243300	F	M00057200B:D04	UC2-PrimColon
2738	648379	1900.K22.gz43_243309	F	M00057211B:A08	UC2-PrimColon
2739	649490	1900.M22.gz43_243311	F	M00057214B:B12	UC2-PrimColon
2740	186522	1900.N22.gz43_243312	F	M00057216C:D12	UC2-PrimColon
2741	561626	1900.E24.gz43_243335	F	M00057204C:G06	UC2-PrimColon
2742	653616	1900.K24.gz43_243341	F	M00057211B:C09	UC2-PrimColon
2743	734828	1909.G01.gz43_243353	F	M00057225C:H07	UC2-PrimColon
2744	427113	1909.H01.gz43 243354	F	M00057226C:E05	UC2-PrimColon
2745	561877	1909.K01.gz43 243357	F	M00057231B:A01	UC2-PrimColon
2746	454563	1909.M01.gz43 243359	F	M00057233B:G04	UC2-PrimColon
2747	625810	1909.P01.gz43 243362	F	M00057236D:H09	UC2-PrimColon
2748	475562	1909.F02.gz43_243368	F	M00057224C:B02	UC2-PrimColon
2749	735749	1909.K02.gz43 243373	F	M00057231C:B04	UC2-PrimColon
2750	289328	1909.B03.gz43 243380	F	M00057219D:G11	UC2-PrimColon
2751	465697	1909.C03.gz43 243381	F	M00057220D:E06	UC2-PrimColon
2752	726892	1909.E03.gz43 243383	F	M00057223B:G01	UC2-PrimColon
2753	723959	1909.J03.gz43 243388	F	M00057230C:C05	UC2-PrimColon
2754	733874	1909.N03.gz43 243392	F	M00057234D:A12	UC2-PrimColon
2755	454129	1909.A04.gz43_243395	F	M00057218C:F10	UC2-PrimColon
2756	733868	1909.B04.gz43 243396	F	M00057219D:H04	UC2-PrimColon
2757	642940	1909.H04.gz43_243402	F	M00057226C:F12	UC2-PrimColon
2758	398061	1909.D05.gz43 243414	F	M00057222B:A06	UC2-PrimColon
2759	734690	1909.G05.gz43 243417	F	M00057225D:E01	UC2-PrimColon
2760	601051	1909.N05.gz43 243424	F	M00057234D:E04	UC2-PrimColon
2761	727255	1909.E06.gz43 243431	F	M00057234D:E04	UC2-PrimColon
2762	632499	1909.I06.gz43_243435	F	M00057229A:B03	UC2-PrimColon
2763	737010	1909.K06.gz43_243437	F	M00057231C:E06	UC2-PrimColon
2764	728121	1909.L06.gz43 243438	F	M00057231C.E00	UC2-PrimColon
2765	472811	1909.B08.gz43 243460	F	M00057232B:C02	UC2-PrimColon
2766	648140	1909.H08.gz43 243466	F		
2767	726251	1909.A09.gz43_243466	F	M00057226C:H10 M00057219A:D05	UC2-PrimColon
2768	642631				UC2-PrimColon
2769	598087	1909.B09.gz43_243476	F	M00057220A:D09	UC2-PrimColon
		1909.J09.gz43_243484		M00057230C:G12	UC2-PrimColon
2770	398061	1909.K09.gz43_243485	F	M00057231C:F12	UC2-PrimColon
2771 2772	463513	1909.M09.gz43_243487	F	M00057233D:G12	UC2-PrimColon
2//2	81	1909.F10.gz43_243496	F	M00057225A:C08	UC2-PrimColon

Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	Т	CLONE ID	LIBRARY
2773	735050	1909.H10.gz43 243498	F	M00057226D:B03	UC2-PrimColon
2774	735140	1909.K10.gz43 243501	F	M00057231C:G04	UC2-PrimColon
2775	220107	1909.O10.gz43_243505	F	M00057236A;F08	UC2-PrimColon
2776	735817	1909.P10.gz43 243506	F	M00057237B:D10	UC2-PrimColon
2777	735514	1909.D11.gz43 243510	F	M00057222D:C10	UC2-PrimColon
2778	602673	1909.F11.gz43 243512	F	M00057225A:E03	UC2-PrimColon
2779	647815	1909.I11.gz43_243515	F	M00057229B:F11	UC2-PrimColon
2780	402024	1909.A12.gz43 243523	F	M00057219A:E11	UC2-PrimColon
2781	631111	1909.C12.gz43_243525	F	M00057221B:B01	UC2-PrimColon
2782	451888	1909.I12.gz43_243531	F	M00057229B:G11	UC2-PrimColon
2783	456224	1909.J12.gz43_243532	F	M00057230D:C05	UC2-PrimColon
2784	555336	1909.L12.gz43_243534	F	M00057232D:B03	UC2-PrimColon
2785	736988	1909.A13.gz43 243539	F	M00057219A;H11	UC2-PrimColon
2786	613067	1909.B13.gz43 243540	F	M00057220B;A06	UC2-PrimColon
2787	736738	1909.E13.gz43 243543	F	M00057223D:H03	UC2-PrimColon
2788	646552	1909.G13.gz43 243545	F	M00057226A:B04	UC2-PrimColon
2789	725120	1909.O13.gz43 243553	F	M00057236B:D11	UC2-PrimColon
2790	591979	1909.A14.gz43 243555	F	M00057219B:B10	UC2-PrimColon
2791	731467	1909.N14.gz43 243568	F	M00057235B:A07	UC2-PrimColon
2792	474869	1909.A15.gz43 243571	F	M00057219B:C06	UC2-PrimColon
2793	735306	1909.C15.gz43 243573	F	M00057221B;E11	UC2-PrimColon
2794	732736	1909.K15.gz43_243581	F	M00057231D:A05	UC2-PrimColon
2795	724417	1909.L15.gz43_243582	F	M00057233A:C04	UC2-PrimColon
2796	454355	1909.E16.gz43_243591	F	M00057224A:D07	UC2-PrimColon
2797	732899	1909.D17.gz43_243606	F	M00057223A:F06	UC2-PrimColon
2798	733149	1909.J17.gz43_243612	F	M00057231A:D04	UC2-PrimColon
2799	735622	1909.B18.gz43_243620	F	M00057220B:G10	UC2-PrimColon
2800	453132	1909.O19.gz43_243649	F	M00057236C:C07	UC2-PrimColon
2801	724781	1909.P19.gz43_243650	F	M00057237D:C11	UC2-PrimColon
2802	556852	1909.B20.gz43_243652	F	M00057220C:A08	UC2-PrimColon
2803	732159	1909.H20.gz43_243658	F	M00057227B:A05	UC2-PrimColon
2804	734609	1909.D21.gz43_243670	F	M00057223B;A07	UC2-PrimColon
2805	616985	1909.I21.gz43_243675	F	M00057230B:B07	UC2-PrimColon
2806	734151	1909.N21.gz43_243680	F	M00057235C:C08	UC2-PrimColon
2807	553850	1909.C22.gz43_243685	F	M00057221'C:E07	UC2-PrimColon
2808	736894	1909.D22.gz43_243686	F	M00057223B:B04	UC2-PrimColon
2809	486051	1909.K22.gz43_243693	F	M00057231D:F10	UC2-PrimColon
2810	734553	1909.N22.gz43_243696	F	M00057235C:F03	UC2-PrimColon
2811	473578	1909.P22.gz43_243698	F	M00057237D:D09	UC2-PrimColon
2812	737010	1909.A23.gz43_243699	F	M00057219D:C02	UC2-PrimColon
2813	735514	1909.B23.gz43_243700	F	M00057220C:F08	UC2-PrimColon
2814	418682	1909.C23.gz43_243701	F	M00057221C:F02	UC2-PrimColon

Table 2

	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
2815	734562	1909.F23.gz43 243704	F	M00057225C:F09	UC2-PrimColon
2816	664711	1909.M23.gz43 243711	F	M00057234C:D11	UC2-PrimColon
2817	727558	1909.E24.gz43 243719	F	M00057224B;H02	UC2-PrimColon
2818	730028	1910.C01.gz43 243733	F	M00057240A:F03	UC2-PrimColon
2819	611604	1910.E01.gz43 243735	F	M00057242C:G12	UC2-PrimColon
2820	558617	1910.H01.gz43_243738	F	M00057246A:G11	UC2-PrimColon
2821	733144	1910.J01.gz43_243740	F	M00057248B:D11	UC2-PrimColon
2822	647242	1910.M01.gz43_243743	F	M00057252B:E04	UC2-PrimColon
2823	733552	1910.F02.gz43_243752	F	M00057243C:D01	UC2-PrimColon
2824	729428	1910.N02.gz43_243760	F	M00057253C:A06	UC2-PrimColon
2825	641680	1910.A03.gz43_243763	F	M00057238A:D07	UC2-PrimColon
2826	514838	1910.L03.gz43_243774	F	M00057251A:F02	UC2-PrimColon
2827	735403	1910.N03.gz43_243776	F	M00057253C:D11	UC2-PrimColon
2828	732859	1910.I04.gz43_243787	F	M00057247C:C11	UC2-PrimColon
2829	730046	1910.C06.gz43_243813	F	M00057240B:C01	UC2-PrimColon
2830	650067	1910.E06.gz43 243815	F	M00057242D:C07	UC2-PrimColon
2831	734884	1910.F06.gz43_243816	F	M00057243C:H11	UC2-PrimColon
2832	550730	1910.I06.gz43_243819	F	M00057247C:F10	UC2-PrimColon
2833	449042	1910.K06,gz43 243821	F	M00057249D:H09	UC2-PrimColon
2834	647704	1910.B07.gz43_243828	F	M00057239B:F05	UC2-PrimColon
2835	555641	1910.G07.gz43_243833	F	M00057245A:F03	UC2-PrimColon
2836	734226	1910.D08.gz43_243846	F	M00057241D:C04	UC2-PrimColon
2837	551437	1910.A09.gz43 243859	F	M00057238B:F05	UC2-PrimColon
2838	639494	1910.M09.gz43_243871	F	M00057252D:B10	UC2-PrimColon
2839	661194	1910.P09.gz43_243874	F	M00057256B:A05	UC2-PrimColon
2840	734466	1910.B10.gz43_243876	F	M00057239C:E05	UC2-PrimColon
2841	734371	1910.I10.gz43_243883	F	M00057247D:D10	UC2-PrimColon
2842	731392	1910.K11.gz43_243901	F	M00057250B:A03	UC2-PrimColon
2843	484964	1910.G12.gz43_243913	F	M00057245B:E02	UC2-PrimColon
2844	733991	1910.H12.gz43_243914	F	M00057246C:B12	UC2-PrimColon
2845	585976	1910.M12.gz43_243919	F	M00057252D:D04	UC2-PrimColon
2846	729125	1910.B13.gz43_243924	F	M00057239C:G07	UC2-PrimColon
2847	733855	1910.C13.gz43_243925	F	M00057240C:A06	UC2-PrimColon
2848	733945	1910.D14.gz43_243942	F	M00057241D:G01	UC2-PrimColon
2849	555103	1910.J14.gz43_243948	F	M00057249A:C06	UC2-PrimColon
2850	726011	1910.K14.gz43_243949	F	M00057250B:D04	UC2-PrimColon
2851	557353	1910.N14.gz43_243952	F	M00057254A:G07	UC2-PrimColon
2852	457846	1910.E15.gz43_243959	F	M00057243A:H03	UC2-PrimColon
2853	456753	1910.B16.gz43_243972	F	M00057239D:D01	UC2-PrimColon
2854	731238	1910,M17.gz43_243999	F	M00057253A:C05	UC2-PrimColon
2855	558332	1910.017.gz43_244001	F	M00057255C:A07	UC2-PrimColon
2856	649390	1910.P17.gz43_244002	F	M00057256D:A11	UC2-PrimColon
2856	649390	1910.P17.gz43_244002	F	M00057256D:A11	UC2-PrimColon

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	able 2				
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SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	LIBRARY
2857	734808	1910.D19.gz43_244022	F	M00057242A:H11	UC2-PrimColon
2858	731125	1910.L19.gz43_244030	F	M00057252A:D10	UC2-PrimColon
2859	476455	1910.G20.gz43_244041	F	M00057245D:G02	UC2-PrimColon
2860	735754	1910.M20.gz43_244047	F	M00057253A:H10	UC2-PrimColon
2861	735477	1910.D21.gz43_244054	F	M00057242B:F07	UC2-PrimColon
2862	734894	1910.I21.gz43_244059	F	M00057248A:H10	UC2-PrimColon
2863	473588	1910.J21.gz43_244060	F	M00057249C:C07	UC2-PrimColon
2864	735665	1910.K21.gz43_244061	F	M00057250C:G02	UC2-PrimColon
2865	725095	1910.M22.gz43_244079	F	M00057253B:C06	UC2-PrimColon
2866	726081	1910.P23.gz43_244098	F	M00057257A:H10	UC2-PrimColon
2867	455821	1910.F24.gz43 244104	F	M00057244C:E06	UC2-PrimColon
2868	449537	1910.024.gz43 244113	F	M00057255D:E02	UC2-PrimColon
2869	514697	1910.P24.gz43 244114	F	M00057257B:C11	UC2-PrimColon
2870	586794	1911.I01.gz43 244123	F	M00057269D:F02	UC2-PrimColon
2871	648710	1911.M01.gz43 244127	F	M00057275B:B02	UC2-PrimColon
2872	630348	1911.C02.gz43 244133	F	M00057260A:E05	UC2-PrimColon
2873	733570	1911.F02.gz43 244136	F	M00057266D:B12	UC2-PrimColon
2874	480723	1911.J02.gz43 244140	F	M00057271A:E04	UC2-PrimColon
2875	728756	1911.C03.gz43 244149	F	M00057260A:E11	UC2-PrimColon
2876	735071	1911.E04.gz43 244167	F	M00057265D:B12	UC2-PrimColon
2877	630655	1911.K04.gz43 244173	F	M00057273A:C08	UC2-PrimColon
2878	736014	1911.N04.gz43 244176	F	M00057279A:G02	UC2-PrimColon
2879	42	1911.P04,gz43 244178	F	M00057283A:E06	UC2-PrimColon
2880	482145	1911.D07.gz43_244214	F	M00057262B;C03	UC2-PrimColon
2881	735834	1911.K10.gz43 244269	F	M00057273B:A12	UC2-PrimColon
2882	42	1911.P10.gz43 244274	F	M00057283B:D09	UC2-PrimColon
2883	736014	1911.M11.gz43 244287	F	M00057277B;C09	UC2-PrimColon
2884	450692	1911.011.gz43 244289	F	M00057281A:D08	UC2-PrimColon
2885	4470	1911.M12.gz43 244303	F	M00057277B;E10	UC2-PrimColon
2886	456065	1911.012.gz43 244305	F	M00057281A:H02	UC2-PrimColon
2887	482985	1911.K15.gz43 244349	F	M00057281A:1102	UC2-PrimColon
2888	652782	1911.G18.gz43 244393	F	M00057268B:B03	UC2-PrimColon
2889	725784	1911.H18.gz43 244394	F	M00057269C:E01	UC2-PrimColon
2890	729321	1911.K19.gz43 244413	F	M00037209C.E01	UC2-PrimColon
2891	732213		F	M00057273D:A06	UC2-PrimColon
		1911.H22.gz43_244458	_		
2892	420504	1911.B23.gz43_244468	F	M00057259D:D11	UC2-PrimColon
2893	735687	1911.F23.gz43_244472	F	M00057267C:G09	UC2-PrimColon
2894	728790	1911.J23.gz43_244476	F	M00057272C:D08	UC2-PrimColon
2895	734131	1911.P23.gz43_244482	F	M00057283D:B12	UC2-PrimColon
2896	725951	1911.D24.gz43_244486	F	M00057265C:F03	UC2-PrimColon
2897	554703	1911.E24.gz43_244487	F	M00057266C:G12	UC2-PrimColon
2898	546642	1911.L24.gz43_244494	F	M00057275B:A12	UC2-PrimColon

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	able 2				
SEQ			ORIEN		
ID NO	CLUSTER	SEQ NAME	T	CLONE ID	_LIBRARY
2899	730189	1912.E03.gz43_244535	F	M00057291B:H08	UC2-PrimColon
2900	728768	1912.D08.gz43_244614	F	M00057290B:A02	UC2-PrimColon
2901	420402	1912.H10.gz43_244650	F	M00057299C;A08	UC2-PrimColon
2902	737109	1912.F12.gz43_244680	F	M00057293A:H03	UC2-PrimColon
2903	424672	1912.H15.gz43_244730	F	M00057299D:E04	UC2-PrimColon
2904	558045	1912.D16.gz43_244742	F	M00057290D:G03	UC2-PrimColon
2905	733856	1912.F16.gz43_244744	F	M00057293B:H04	UC2-PrimColon
2906	736449	1912.H17.gz43_244762	F	M00057300A:A08	UC2-PrimColon
2907	148201	1912.H18.gz43_244778	F	M00057300A:B06	UC2-PrimColon
2908	639427	1912.E19.gz43_244791	F	M00057292A:B08	UC2-PrimColon
2909	730187	1912.B21.gz43_244820	F	M00057287A:H06	UC2-PrimColon
2910	732300	1912.D24.gz43_244870	F	M00057291B:D08	UC2-PrimColon
2911	465528	1921.A01.gz43_244883	F	M00057314A:A10	UC2-PrimColon
2912	549552	1921.J01.gz43_244892	F	M00057331D:D08	UC2-PrimColon
2913	470199	1921.E02.gz43_244903	F	M00057320A:G08	UC2-PrimColon
2914	729125	1921.F03.gz43_244920	F	M00057323B:G04	UC2-PrimColon
2915	640341	1921.C04.gz43_244933	F	M00057316D:D02	UC2-PrimColon
2916	77737	1921.F04.gz43_244936	F	M00057323B:H08	UC2-PrimColon
2917	735469	1921.G04.gz43_244937	F	M00057324B:E04	UC2-PrimColon
2918	736733	1921.B06.gz43_244964	F	M00057315D:C06	UC2-PrimColon
2919	732969	1921.I06.gz43_244971	F	M00057328C:F11	UC2-PrimColon
2920	635849	1921.J06.gz43_244972	F	M00057332A;C06	UC2-PrimColon
2921	733146	1921.006.gz43_244977	F	M00057339D:H09	UC2-PrimColon
2922	737114	1921.A07.gz43_244979	F	M00057314B:H06	UC2-PrimColon
2923	732114	1921.L07.gz43_244990	F	M00057334D:E03	UC2-PrimColon
2924	482788	1921.P08.gz43 245010	F	M00057341D:B09	UC2-PrimColon
2925	554647	1921.M10.gz43_245039	F	M00057337C:G12	UC2-PrimColon
2926	552623	1921.B11.gz43_245044	F	M00057316A:B04	UC2-PrimColon
2927	447705	1921.E11.gz43_245047	F	M00057320D:C02	UC2-PrimColon
2928	731748	1921.G11.gz43_245049	F	M00057324C:G05	UC2-PrimColon
2929	727093	1921.B12.gz43 245060	F	M00057316A:D09	UC2-PrimColon
2930	551607	1921.N12.gz43_245072	F	M00057339A;E08	UC2-PrimColon
2931	727013	1921.G13.gz43_245081	F	M00057324D:E10	UC2-PrimColon
2932	727018	1921.L13.gz43 245086	F	M00057336B;E01	UC2-PrimColon
2933	548635	1921.G14.gz43 245097	F	M00057324D:H03	UC2-PrimColon
2934	726692	1921.J14.gz43_245100	F	M00057332C:F12	UC2-PrimColon
2935	556430	1921.D15.gz43_245110	F	M00057319D:E06	UC2-PrimColon
2936	726692	1921.A16.gz43 245123	F	M00057314D:E09	UC2-PrimColon
2937	473225	1921.F16.gz43_245128	F	M00057323D:E10	UC2-PrimColon
2938	892	1921.K16.gz43_245133	F	M00057334A:C12	UC2-PrimColon
2939	730997	1921.C17.gz43_245141	F	M00057318B:H05	UC2-PrimColon
2940	51616	1921.E18.gz43_245159	F	M00057323A:F01	UC2-PrimColon

Table 2

SEQ ORIE NT CLONE ID LIBRAN	Colon
NO CLUSTER	Colon
2941 \$46642 1921.F18.gz43 245160 F M00057324A:A09 UC2-Primt	Colon
2942 643005 1921.G18.gz43 245161 F M00057325B:D06 UC2-Primt	Colon
2944 735054 1921.H18.grd3 245162 F M00057326D.B10 UC2-Print	Colon
Page	Colon
2945 733464 1921,J19,gr43 245180 F M0005733A:D08 UC2-Primt 2946 457092 1921,B20,gr43 245188 F M00057316C:A06 UC2-Primt 2947 733723 1921,H21,gr43 245211 F M00057331C:E10 UC2-Primt 2948 557177 1921,I21,gr43 245211 F M00057331C:E10 UC2-Primt 2950 448431 1921,L21,gr43 245214 F M00057336D:F07 UC2-Primt 2951 697006 1921,L22,gr43 245226 F M00057336D:F07 UC2-Primt 2952 452936 1921,L22,gr43 245227 F M00057331C:F10 UC2-Primt 2953 727132 1921,L22,gr43 245230 F M00057331C:F10 UC2-Primt 2954 513306 1921,F23,gr43 245230 F M00057334B:A06 UC2-Primt 2955 642986 1921,P24,gr43 245266 F M00057334A:G07 UC2-Primt 2957 498509 <t< td=""><td>Colon Colon Colon Colon Colon Colon Colon Colon Colon Colon Colon</td></t<>	Colon
2946 457092 1921.B20.gz43 245188 F M00057316C:A06 UC2-Print	Colon Colon Colon Colon Colon Colon Colon Colon Colon
2944 733723 1921.H21_gr43 245210 F M00057328A:F02 UC2-Primt	Colon Colon Colon Colon Colon Colon
2948 557177 1921.121_gz43 245211 F M00057331C.E10 UC2-Primt	Colon Colon Colon Colon Colon
2949 734582 1921.K21.gz43 245213 F M00057334B:F01 UC2-Primt	Colon Colon Colon Colon
2950 448431 1921.L21_gzd3_245214 F M00057336D:F07 UC2-Primt	Colon Colon Colon
2951 697006 1921.H22_gz43 245226 F M00057328A:F10 UC2-Primt	Colon Colon
2952 452936 1921.122_gz43 245227 F M00057331C:F10 UC2-Prim(2953 727132 1921.1.22_gz43 245230 F M00057331A:A06 UC2-Prim(2954 513306 1921.P32_gz43 245240 F M00057334R:A06 UC2-Prim(2955 642986 1921.P24_gz43 245266 F M0005734AR:AG07 UC2-Prim(2956 496752 1924.G03_gz43 245338 F M0004330A:D09 UC2-ColomM 2957 498509 1924.H03_gz43 245333 F M0004330A:D09 UC2-ColomM 2958 450805 1924.E04_gz43 245352 F M0004330A:F06 UC2-ColomM 2959 451456 1924.G04_gz43 245354 F M0004330B:F06 UC2-ColomM 2950 494625 1924.M04_gz43 245356 F M0004330B:F06 UC2-ColomM 2950 494625 1924.M04_gz43 245356 F M0004331B:E10 UC2-ColomM 2961 490401 1924.P05_gz43 245379 F M00043321B:E05 UC2-ColomM 2961 490401 1924.P05_gz43 245379 F M0004321B:E05 UC2-ColomM 2961 490401 1924.P05_gz43 245379 F M00043204.P06 UC2-ColomM 2961 490401 1924.P05_gz43	Colon
2953 727132 1921.L22 gz43 245230 F M00057337A:A06 UC2-Primt 2954 513306 1921.P23 gz43 245240 F M00057324B:A06 UC2-Primt 2955 642986 1921.P24 gz43 245266 F M00057344A:G07 UC2-Primt 2956 496752 1924.G03 gz43 245338 F M00043308A:D09 UC2-ColomM 2957 498509 1924.H03 gz43 245339 F M00043310A:F01 UC2-ColomM 2958 450805 1924.G04 gz43 245352 F M00043308A:D0 UC2-ColomM 2959 451456 1924.G04 gz43 245354 F M00043308A:F06 UC2-ColomM 2960 494625 1924.M04 gz43 245369 F M00043317B:B12 UC2-ColomM 2961 490401 1924.P05 gz43 245379 F M0004331B:E05 UC2-ColomM 2961 490401 1924.P05 gz43 245379 F M0004331B:E05 UC2-ColomM	
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2954 513306 1921.F23.gz43 245240 F M00057324B:A06 UC2-PrimC 2955 642986 1921.P24.gz43 245266 F M00057324A:G07 UC2-PrimC 2956 496752 1924.G03.gz43 245333 F M00043308A:D09 UC2-ColonM 2957 498509 1924.H03.gz43 245339 F M00043310A:F01 UC2-ColonM 2958 450805 1924.E04.gz43 245352 F M0004330B:A10 UC2-ColonM 2950 494625 1924.G04.gz43 245356 F M0004331B:B31 UC2-ColonM 2960 494625 1924.M04.gz43 245369 F M0004331B:E05 UC2-ColonM 2961 490401 1924.P05.gz43 245379 F M0004331B:E05 UC2-ColonM	_oion
2956 496752 1924.G03.gz43 245338 F M00043308A:D09 UC2-ColonM 2957 498509 1924.H03.gz43 245339 F M00043310A:F01 UC2-ColonM 2958 450805 1924.E04.gz43 245352 F M00043304B:A10 UC2-ColonM 2959 451456 1924.G04.gz43 245354 F M00043308A:F06 UC2-ColonM 2960 494625 1924.M04.gz43 245360 F M00043317B:B12 UC2-ColonM 2961 490401 1924.P05.gz43 245379 F M00043321B:E05 UC2-ColonM	
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2958 450805 1924.E04.gz43 245352 F M00043304B:A10 UC2-ColonM 2959 451456 1924.G04.gz43 245354 F M00043308A:F06 UC2-ColonM 2960 494625 1924.M04.gz43 245360 F M00043317B:B12 UC2-ColonM 2961 490401 1924.P05.gz43 245379 F M00043321B:E05 UC2-ColonM	etLiver
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2960 494625 1924.M04.gz43_245360 F M00043317B:B12 UC2-ColonM 2961 490401 1924.P05.gz43_245379 F M00043321B:E05 UC2-ColonM	etLiver
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2961 490401 1924.P05.gz43_245379 F M00043321B:E05 UC2-ColonM	ctLiver
	etLiver
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2964 494306 1924.I10.gz43 245452 F M00043312D:A02 UC2-ColonM	
2965 492544 1924.J11.gz43_245469 F M00043313D:B04 UC2-ColonM	etLiver
2966 451081 1924.M11.gz43 245472 F M00043317D:C02 UC2-ColonM	etLiver
2967 495951 1924.K14.gz43 245518 F M00043315C:D05 UC2-ColonM	etLiver
2968 450349 1924.D15.gz43 245527 F M00043303B:E11 UC2-ColonM	etLiver
2969 498951 1924.L15.gz43_245535 F M00043316C:F06 UC2-ColonM	etLiver
2970 450999 1924.C17.gz43 245558 F M00043301D:B12 UC2-ColonM	etLiver
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2972 498242 1924.F18.gz43 245577 F M00043307B:F11 UC2-ColonM	
2973 499700 1924.H18.gz43 245579 F M00043311B:H08 UC2-ColonM	
2974 479604 1924.L19.gz43 245599 F M00043316D:F04 UC2-ColonM	etLiver
2975 562516 1924.N19.gz43 245601 F M00043319A;D01 UC2-ColonM	
2976 553779 1924.L21.gz43 245631 F M00043316D:F09 UC2-ColonM	
2977 493746 1924.E24.gz43 245672 F M00043306B:A09 UC2-ColonM	
2978 494325 1924.H24.gz43 245675 F M00043312B;A10 UC2-ColonM	
2979 453078 1933.H03.gz43 245723 F M00043340B:B04 UC2-ColonM	
2980 492887 1933.J04.gz43 245741 F M00043343D:H03 UC2-ColonM	etLiver
2981 450283 1933.F05.gz43 245753 F M00043335D:E02 UC2-ColonM	
2982 494271 1933.M05.gz43 245760 F M00043351C:A07 UC2-ColonM	etLiver

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Table 2

	able 2				
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2984	412621	1933.D06.gz43_245767	F	M00043330D:G05	UC2-ColonMetLiver
2985	492629	1933.I06.gz43_245772	F	M00043342C:G02	UC2-ColonMetLiver
2986	491212	1933.E07.gz43_245784	F	M00043334A:F10	UC2-ColonMetLiver
2987	447057	1933.H09.gz43_245819	F	M00043340D:C03	UC2-ColonMetLiver
2988	496909	1933.C10.gz43_245830	F	M00043328A:E12	UC2-ColonMctLiver
2989	491933	1933.D10.gz43_245831	F	M00043331C:G01	UC2-ColonMetLiver
2990	496870	1933.N10.gz43_245841	F	M00043353D:E12	UC2-ColonMetLiver
2991	500040	1933.B12.gz43_245861	F	M00043326A:H06	UC2-ColonMetLiver
2992	553594	1933.O12.gz43_245874	F	M00043354D:H08	UC2-ColonMetLiver
2993	561712	1933.N14.gz43_245905	F	M00043354A:E06	UC2-ColonMetLiver
2994	450027	1933.J15.gz43 245917	F	M00043344C:C11	UC2-ColonMetLiver
2995	450129	1933,K16,gz43_245934	F	M00043348A:D02	UC2-ColonMetLiver
2996	492779	1933.C17.gz43_245942	F	M00043329B:H07	UC2-ColonMetLiver
2997	453068	1933,C18,gz43_245958	F	M00043329C:D12	UC2-ColonMetLiver
2998	450804	1933.F18.gz43_245961	F	M00043336C:A04	UC2-ColonMetLiver
2999	1015	1933.N18.gz43 245969	F	M00043354B:C04	UC2-ColonMetLiver
3000	561834	1933.N20.gz43 246001	F	M00043354B:F12	UC2-ColonMetLiver
3001	552687	1933,O23,gz43 246050	F	M00043355B:B01	UC2-ColonMetLiver
3002	517346	1933.O24.gz43 246066	F	M00043355B:D12	UC2-ColonMetLiver
3003	491448	1935.B05.gz43 246289	F	M00043392C:F02	UC2-ColonMetLiver
3004	553890	1935,K06,gz43 246314	F	M00043501A:D07	UC2-ColonMetLiver
3005	635439	1935.N06.gz43_246317	F	M00043504B:C02	UC2-ColonMetLiver
3006	402070	1935.I07.gz43 246328	F	M00043409C;C07	UC2-ColonMetLiver
3007	448924	1935.H08.gz43 246343	F	M00043406D;C04	UC2-ColonMetLiver
3008	562886	1935.O08.gz43 246350	F	M00043505B:G03	UC2-ColonMetLiver
3009	558055	1935.O09.gz43 246366	F	M00043505B:G07	UC2-ColonMetLiver
3010	451456	1935.H15.gz43 246455	F	M00043407C:H08	UC2-ColonMetLiver
3011	555399	1935.N16.gz43 246477	F	M00043504C:E03	UC2-ColonMetLiver
3012	498629	1935,H17,gz43 246487	F	M00043407D:G06	UC2-ColonMetLiver
3013	490805	1935.E18.gz43 246500	F	M00043401A:E09	UC2-ColonMetLiver
3014	450335	1935.F18.gz43 246501	F	M00043403B:A12	UC2-ColonMetLiver
3015	446964	1935.N22.gz43 246573	F	M00043504D:G08	UC2-ColonMetLiver
3016	522703	1935.P24.gz43 246607	F	M00043506D;F06	UC2-ColonMetLiver
3017	549853	1936,K05,gz43 246682	F	M00054494A:H04	UC2-ColonMetLiver
3018	560275	1936,M05,gz43 246684	F	M00054497B:C10	UC2-ColonMetLiver
3019	481057	1936.P05.gz43 246687	F	M00054501C:E08	UC2-ColonMetLiver
3020	414739	1936.C08.gz43 246722	F	M00043508D:C01	UC2-ColonMetLiver
3021	559720	1936.O08.gz43 246734	F	M00054500A;F04	UC2-ColonMetLiver
3022	554908	1936,M11,gz43 246780	F	M00054497D;A04	UC2-ColonMetLiver
3023	449000	1936.N11.gz43 246781	F	M00054498D:F01	UC2-ColonMetLiver
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Table 2

1	able 2				
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3026	489368	1936.J16.gz43_246857	F	M00054493C:E04	UC2-ColonMetLiver
3027	562823	1936.B18.gz43_246881	F	M00043508B:G11	UC2-ColonMetLiver
3028	559574	1936.E21.gz43_246932	F	M00054487C:A01	UC2-ColonMetLiver
3029	552920	1936.O23.gz43_246974	F	M00054501A:E11	UC2-ColonMetLiver
3030	552086	1971.A01.gz43_246992	F	M00054717A:A03	UC2-ColonMetLiver
3031	553123	1971.D01.gz43_246995	F	M00054720C:A01	UC2-ColonMetLiver
3032	561636	1971.H01.gz43_246999	F	M00054724D:C05	UC2-ColonMetLiver
3033	496772	1971.K01.gz43_247002	F	M00054727D:C06	UC2-ColonMetLiver
3034	549550	1971.A03.gz43_247024	F	M00054717A:C07	UC2-ColonMetLiver
3035	555837	1971.B05.gz43_247057	F	M00054718B:D03	UC2-ColonMetLiver
3036	557714	1971.D05.gz43_247059	F	M00054720C:G10	UC2-ColonMetLiver
3037	446999	1971.E05.gz43_247060	F	M00054721C:D11	UC2-ColonMetLiver
3038	450410	1971.F05.gz43_247061	F	M00054722C:D01	UC2-ColonMetLiver
3039	484043	1971.L05.gz43_247067	F	M00054729B:A08	UC2-ColonMetLiver
3040	452488	1971.N05.gz43_247069	F	M00054731A:D07	UC2-ColonMetLiver
3041	450278	1971.A06.gz43_247072	F	M00054717A:G09	UC2-ColonMetLiver
3042	561313	1971.D06.gz43_247075	F	M00054720C:H01	UC2-ColonMetLiver
3043	557426	1971.L06.gz43_247083	F	M00054729B;B03	UC2-ColonMetLiver
3044	553316	1971.F09.gz43_247125	F	M00054722D:C08	UC2-ColonMetLiver
3045	555818	1971.I09.gz43_247128	F	M00054726A:D02	UC2-ColonMetLiver
3046	551096	1971.H10.gz43_247143	F	M00054725A:A05	UC2-ColonMetLiver
3047	556654	1971.P10.gz43_247151	F	M00054734B;C06	UC2-ColonMetLiver
3048	555491	1971.H11.gz43_247159	F	M00054725A:F09	UC2-ColonMetLiver
3049	559380	1971.M11.gz43_247164	F	M00054730B:F11	UC2-ColonMetLiver
3050	86175	1971.H12.gz43_247175	F	M00054725A:H06	UC2-ColonMetLiver
3051	561830	1971.L12.gz43_247179	F	M00054729C:C08	UC2-ColonMetLiver
3052	421959	1971.G14.gz43_247206	F	M00054724A:G02	UC2-ColonMetLiver
3053	448453	1971.N14.gz43_247213	F	M00054731B:D04	UC2-ColonMetLiver
3054	562263	1971.L15.gz43_247227	F	M00054729C:G07	UC2-ColonMetLiver
3055	553739	1971.I16.gz43_247240	F	M00054726B:F08	UC2-ColonMetLiver
3056	557747	1971.117.gz43_247256	F	M00054726B:F09	UC2-ColonMetLiver
3057	14573	1971.N17.gz43_247261	F	M00054731B:G02	UC2-ColonMetLiver
3058	553108	1971.P17.gz43_247263	F	M00054734C;A07	UC2-ColonMetLiver
3059	554048	1971.K18.gz43_247274	F	M00054728D:B07	UC2-ColonMetLiver
3060	550315	1971.L18.gz43_247275	F	M00054729D:D05	UC2-ColonMetLiver
3061	449035	1971.I21.gz43_247320	F	M00054726C:D10	UC2-ColonMetLiver
3062	524721	1971.I22.gz43_247336	F	M00054726C:E07	UC2-ColonMetLiver
3063	555571	1971.A24.gz43_247360	F	M00054718A:D11	UC2-ColonMetLiver
3064	553869	1971,N24.gz43_247373	F	M00054731C:H12	UC2-ColonMetLiver
3065	552686	1971.P24.gz43_247375	F	M00054735A:G09	UC2-ColonMetLiver
3066	424723	1972.103.gz43_247416	F	M00054743A:C07	UC2-ColonMetLiver

WO 02/14500 Table 2

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Section 1972,P03,g243 247423 F M00054750D:H12 UC2-ColomMetLiver 3068 S1441 1972,L04,g243 247435 F M00054751A:A05 UC2-ColomMetLiver 3070 S56167 1972,C05,g243 247442 F M00054751A:A05 UC2-ColomMetLiver 3071 S53834 1972,J05,g243 247442 F M0005473B:H11 UC2-ColomMetLiver 3071 S53834 1972,J05,g243 247442 F M0005473B:H11 UC2-ColomMetLiver 3073 466235 1972,E08,g243 247442 F M000547376:B01 UC2-ColomMetLiver 3073 466235 1972,E08,g243 247484 F M00054739C:B12 UC2-ColomMetLiver 3074 562323 1972,009,g243 247518 F M00054739C:B12 UC2-ColomMetLiver 3075 45042 1972,E112,g243 247565 F M00054739C:B12 UC2-ColomMetLiver 3076 557568 1972,012,g243 247566 F M00054739C:B12 UC2-ColomMetLiver 3077 522322 1972,B13,g243 247567 F M00054736E:B10 UC2-ColomMetLiver 3078 52576 972,D13,g243 247571 F M00054736E:B10 UC2-ColomMetLiver 3079 452257 1972,I13,g243 247571 F M00054736E:B10 UC2-ColomMetLiver 3080 523753 1972,N13,g243 247571 F M00054736E:B10 UC2-ColomMetLiver 3081 5920 1972,N14,g243 247597 F M00054748E:B10 UC2-ColomMetLiver 3082 553702 1972,P14,g243 247597 F M00054748E:B10 UC2-ColomMetLiver 3084 49312 1972,D15,g243 247603 F M00054748E:B10 UC2-ColomMetLiver 3084 49312 1972,D15,g243 247605 F M0005474E:B12 UC2-ColomMetLiver 3085 562793 1972,R15,g243 247605 F M0005474C:B12 UC2-ColomMetLiver 3085 55378 1972,G15,g243 247605 F M0005474C:B12 UC2-ColomMetLiver 3085 554793 1972,R15,g243 247605 F M0005474C:B12 UC2-ColomMetLiver 3085 55577 1972,D13,g243 247605 F M0005474C:B12 UC2-ColomMetLiver 3085 55577 1972,D13,g243 247605 F M0005474C:B12 UC2-ColomMetLiver 3085 55577 1972,D13,g243 247605 F M0005474C:B08 UC2-ColomMetLiver 3095 555077 1972,L19,g243 247605 F M0005474C:B08 UC2-ColomMetLiver 3095 555071 1972,						
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3071 553834 1972.105.gz43 247449 F M0005474A.G10 UC2-ColonMetLiver 3072 454186 1972.C06.gz43 247458 F M00054739C:B10 UC2-ColonMetLiver 3073 466235 1972.C08.gz43 247452 F M00054739C:B10 UC2-ColonMetLiver 3074 562323 1972.009.gz43 247518 F M00054739C:B10 UC2-ColonMetLiver 3075 450142 1972.E12.gz43 247556 F M00054739C:E05 UC2-ColonMetLiver 3076 557568 1972.D12.gz43 247566 F M00054739C:E05 UC2-ColonMetLiver 3077 52232 1972.B13.gz43 247566 F M00054736B:H03 UC2-ColonMetLiver 3078 558679 1972.D13.gz43 247576 F M00054736B:H03 UC2-ColonMetLiver 3079 45227 1972.D13.gz43 247576 F M00054736B:H03 UC2-ColonMetLiver 3080 523753 1972.N13.gz43 247567 F M00054748B:G10 UC2-ColonMetLiver 3081 59202 1972.N14.gz43 247597 F M00054748B:G10 UC2-ColonMetLiver 3082 553702 1972.N14.gz43 247599 F M00054748B:G10 UC2-ColonMetLiver 3083 495832 1972.D15.gz43 247603 F M00054748B:G10 UC2-ColonMetLiver 3084 493122 1972.F15.gz43 247605 F M0005474B:D607 UC2-ColonMetLiver 3085 493135 1972.G15.gz43 247605 F M00054740C:D08 UC2-ColonMetLiver 3086 562793 1972.K15.gz43 247605 F M00054740C:D08 UC2-ColonMetLiver 3087 87884 1972.D16.gz43 247605 F M00054740C:D08 UC2-ColonMetLiver 3088 53457 1972.G18.gz43 247605 F M00054740C:D08 UC2-ColonMetLiver 3089 551778 1972.K15.gz43 247605 F M00054740C:D02 UC2-ColonMetLiver 3090 555077 1972.N19.gz43 247670 F M00054745C:D10 UC2-ColonMetLiver 3091 575871 1972.K19.gz43 247670 F M00054745C:D10 UC2-ColonMetLiver 3092 50282 1972.C2.gz43 247698 F M00054745C:D02 UC2-ColonMetLiver 3093 53452 1972.L2.gz43 247767 F M00054745C:D02 UC2-ColonMetLiver 3094 555073 1972.M2.gz43 247767 F M00054745C:D02 UC2-ColonMetLiver 3095 503452 1978.L03.gz43 247767 F M00054765D:D05 UC2-ColonMetLiver 3096	3069	86145	1972.P04.gz43_247439	F	M00054751A:A05	UC2-ColonMetLiver
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3081 59202 1972.N14.gz43 247597 F M00054748B:H09 UC2-ColonMetLiver 3082 553702 1972.P14.gz43 247599 F M00054751B:F12 UC2-ColonMetLiver 3083 495832 1972.D15.gz43 247605 F M0005473B:G07 UC2-ColonMetLiver 3084 493122 1972.F15.gz43 247605 F M0005474C:D08 UC2-ColonMetLiver 3085 562793 1972.K15.gz43 247606 F M0005474C:D12 UC2-ColonMetLiver 3088 553457 1972.O18.gz43 247607 F M0005474C:D20 UC2-ColonMetLiver 3088 553457 1972.O18.gz43 247607 F M0005474C:D02 UC2-ColonMetLiver 3080 551778 1972.O18.gz43 247670 F M0005474D:D02 UC2-ColonMetLiver 3091 555077 1972.L19.gz43 247671 F M00054743D:P04 UC2-ColonMetLiver 3092 56028 1972.C12.gz43 247678 F M00054733B:E02 UC2-ColonMetLiver	3079	452257	1972.I13.gz43_247576	F	M00054743C:E11	UC2-ColonMetLiver
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3083 495832 1972.D15_g243 247603 F M00054738D:G07 UC2-ColonMetLiver 3084 493122 1972.F15_g243 247605 F M00054740C:D18 UC2-ColonMetLiver 3085 493153 1972.G15_g243 247605 F M00054741C:D08 UC2-ColonMetLiver 3086 562793 1972.K15_g243 247610 F M00054745C:B12 UC2-ColonMetLiver 3087 58784 1972.I16_g243 247625 F M00054740C:D01 UC2-ColonMetLiver 3088 553475 1972.C18_g243 247670 F M00054741D:C05 UC2-ColonMetLiver 3089 551778 1972.E19_g243 247672 F M00054741D:C05 UC2-ColonMetLiver 3091 55787 1972.L19_g243 247674 F M00054743D:F04 UC2-ColonMetLiver 3092 560282 1972.C21_g243 247734 F M00054738B:E12 UC2-ColonMetLiver 3094 555773 1972.D42_g243 247474 F M00054739A:F07 UC2-ColonMetLiver </td <td>3081</td> <td>59202</td> <td></td> <td></td> <td>M00054748B:H09</td> <td>UC2-ColonMetLiver</td>	3081	59202			M00054748B:H09	UC2-ColonMetLiver
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3085 493135 1972.G15.g243 247606 F M00054741C:D08 UC2-ColonMetLiver 3086 562793 1972.K15.g243 247610 F M00054745:B12 UC2-ColonMetLiver 3087 587854 1972.I16.g243 247625 F M00054745:CB12 UC2-ColonMetLiver 3088 553457 1972.O18.g243 247670 F M00054741D:C03 UC2-ColonMetLiver 3089 551778 1972.G19.g243 247672 F M00054743.D:F04 UC2-ColonMetLiver 3091 555077 1972.K19.g243 247674 F M00054743.D:F04 UC2-ColonMetLiver 3092 560282 1972.C21.g243 247678 F M00054745.D:E05 UC2-ColonMetLiver 3094 555773 1972.D24.g243 247767 F M00054746D:E05 UC2-ColonMetLiver 3092 560282 1972.D24.g243 247767 F M00054746D:E05 UC2-ColonMetLiver 3094 74309 1972.M24.g243 247767 F M00054746D:E05 UC2-ColonMetLive	3083	495832	1972.D15.gz43_247603	F	M00054738D:G07	UC2-ColonMetLiver
3086 562793 1972.K15.gz43 247610 F M00054745C:B12 UC2-ColonMetLiver 3087 587854 1972.116.gz43 247625 F M00054744C:D02 UC2-ColonMetLiver 3088 553457 1972.018.gz43 247662 F M00054741D:C05 UC2-ColonMetLiver 3089 551778 1972.G19.gz43 247670 F M00054741D:C05 UC2-ColonMetLiver 3090 555077 1972.K19.gz43 247672 F M00054743D:F04 UC2-ColonMetLiver 3091 557871 1972.K19.gz43 247674 F M00054743D:F04 UC2-ColonMetLiver 3092 560282 1972.C21.gz43 247763 F M0005473B:E12 UC2-ColonMetLiver 3093 55373 1972.D24.gz43 247773 F M00054739.FP01 UC2-ColonMetLiver 3094 55573 1972.D24.gz43 247747 F M00054739.FP01 UC2-ColonMetLiver 3095 474309 1972.M24.gz43 247747 F M00054763D:F01 UC2-ColonMetLiver <td>3084</td> <td>493122</td> <td>1972.F15.gz43_247605</td> <td></td> <td>M00054740C:H08</td> <td>UC2-ColonMetLiver</td>	3084	493122	1972.F15.gz43_247605		M00054740C:H08	UC2-ColonMetLiver
3087 587854 1972.116.gz43 247625 F M00054744C:D02 UC2-ColonMetLiver 3088 553457 1972.018.gz43 247662 F M00054750C:D01 UC2-ColonMetLiver 3089 551778 1972.018.gz43 247670 F M00054741D:C05 UC2-ColonMetLiver 3090 555077 1972.119.gz43 247672 F M00054743D:F04 UC2-ColonMetLiver 3091 560282 1972.C11.gz43 247674 F M00054743C:H02 UC2-ColonMetLiver 3093 560282 1972.C12.gz43 247678 F M00054745D:E05 UC2-ColonMetLiver 3094 555773 1972.L22.gz43 247773 F M00054746D:E05 UC2-ColonMetLiver 3095 474309 1972.M24.gz43 247769 F M000547754D:E05 UC2-ColonMetLiver 3096 474309 1981.101.gz43 247789 F M000547754D:F01 UC2-ColonMetLiver 3097 472196 1981.102.gz43 247784 F M00054754D:F01 UC2-ColonMetLiv	3085	493135	1972.G15.gz43_247606	F	M00054741C:D08	UC2-ColonMetLiver
3088 553457 1972.O18,gz43 247662 F M00054750C:D01 UC2-ColonMetLiver 3089 550778 1972.G19,gz43 247670 F M00054741D:C05 UC2-ColonMetLiver 3090 555077 1972.K19,gz43 247671 F M00054743D:F04 UC2-ColonMetLiver 3091 557871 1972.K19,gz43 247674 F M00054749C:H02 UC2-ColonMetLiver 3092 560282 1972.C21,gz43 247768 F M0005473B:B12 UC2-ColonMetLiver 3093 503452 1972.L22,gz43 247773 F M00054736D:E05 UC2-ColonMetLiver 3094 555773 1972.D24,gz43 247756 F M00054739A:F07 UC2-ColonMetLiver 3096 556019 1981.D1,gz43 247769 F M0005476D:F01 UC2-ColonMetLiver 3097 472196 1981.D0,gz43 247797 F M0005476D:F01 UC2-ColonMetLiver 3099 553158 1981.F03,gz43 247780 F M00054763-B08 UC2-ColonMetLiver	3086	562793	1972.K15.gz43_247610		M00054745C:B12	UC2-ColonMetLiver
3089 551778 1972.G19.g243_247670 F M00054741D:C05 UC2-ColonMetLiver 3090 555077 1972.H19.g243_247672 F M00054743D:F04 UC2-ColonMetLiver 3091 557871 1972.K19.g243_247672 F M0005473B:F04 UC2-ColonMetLiver 3092 560282 1972.C21.g243_247698 F M0005473B:E12 UC2-ColonMetLiver 3093 503452 1972.L22.g243_247723 F M00054739.F02 UC2-ColonMetLiver 3094 55573 1972.D44.g243_247474 F M00054739.F02 UC2-ColonMetLiver 3095 474309 1972.M24.g243_24774 F M00054739.F02 UC2-ColonMetLiver 3096 556019 1981.D1.g243_24776 F M00054764D:F01 UC2-ColonMetLiver 3097 472196 1981.D0.g243_247784 F M00054763D:B05 UC2-ColonMetLiver 3098 513632 1981.D0.g243_24784 F M00054763D:B05 UC2-ColonMetLiver 3100 4986.C2 1981.D0.g243_247816 F M00054766C:B08 UC2-ColonM	3087	587854	1972.J16.gz43_247625		M00054744C:D02	UC2-ColonMetLiver
3090 555077 1972.I19.g243 247672 F M00054743D.F04 UC2-ColonMetLiver 3091 557871 1972.K19.g243 247674 F M00054745C.H02 UC2-ColonMetLiver 3092 56282 1972.C21.g243 247674 F M00054733B.E12 UC2-ColonMetLiver 3093 503452 1972.L22.g243 247723 F M0005474D.E05 UC2-ColonMetLiver 3094 74309 1972.D24.g243 247747 F M0005474D.E05 UC2-ColonMetLiver 3095 474309 1972.D24.g243 247769 F M0005476D.F01 UC2-ColonMetLiver 3096 74309 1981.D01.g243 247789 F M0005476D.F01 UC2-ColonMetLiver 3097 472196 1981.D03.g243 247795 F M0005476D.F01 UC2-ColonMetLiver 3098 513632 1981.D03.g243 247797 F M00054759A.B08 UC2-ColonMetLiver 3100 51693 1981.C04.g243 247810 F M00054759A.B08 UC2-ColonMetLiver	3088	553457	1972.O18.gz43_247662		M00054750C:D01	UC2-ColonMetLiver
3091 557871 1972.K19.gz43 247674 F M00054745C:H02 UC2-ColonMetLiver	3089	551778	1972.G19.gz43_247670	F	M00054741D:C05	UC2-ColonMetLiver
3092 560282 1972.C21.gz43 247698 F M00054738B:E12 UC2-ColonMetLiver 3093 53452 1972.L22.gz43 247723 F M00054746D:E05 UC2-ColonMetLiver 3094 55577 1972.D24.gz43 247474 F M00054739A:F07 UC2-ColonMetLiver 3095 474309 1972.M24.gz43 247769 F M0005476D:F01 UC2-ColonMetLiver 3096 556019 1981.101.gz43 247769 F M00054763D:F01 UC2-ColonMetLiver 3097 472196 1981.103.gz43 247784 F M00054763L:D61 UC2-ColonMetLiver 3098 513632 1981.103.gz43 247797 F M00054759A:B08 UC2-ColonMetLiver 3100 4986.2 1981.103.gz43 247803 F M00054766C:B08 UC2-ColonMetLiver 3101 554021 1981.04.gz43 247810 F M00054769B:D12 UC2-ColonMetLiver 3104 2486.1 1981.04.gz43 247822 F M00054769B:D12 UC2-ColonMetLiver	3090	555077	1972.I19.gz43_247672	F	M00054743D:F04	UC2-ColonMetLiver
3093 503452 1972.122.gz43 247723 F M00054746D:E05 UC2-ColonMetLiver 3094 555773 1972.D24.gz43 247747 F M00054739A:F07 UC2-ColonMetLiver 3095 474309 1972.D24.gz43 247756 F M00054774D:B05 UC2-ColonMetLiver 3096 556019 1981.101.gz43 247769 F M00054764D:F01 UC2-ColonMetLiver 3097 472196 1981.102.gz43 247784 F M00054763A:A10 UC2-ColonMetLiver 3098 513632 1981.D03.gz43 247797 F M00054759A:B08 UC2-ColonMetLiver 3100 498662 1981.L03.gz43 247803 F M00054759A:B08 UC2-ColonMetLiver 3101 551693 1981.C04.gz43 247816 F M00054750-BDF11 UC2-ColonMetLiver 3103 550402 1981.104.gz43 247816 F M00054776B:D02 UC2-ColonMetLiver 3104 448510 1981.P04.gz43 247832 F M00054776B:D10 UC2-ColonMetLive	3091	557871	1972.K19.gz43_247674		M00054745C:H02	UC2-ColonMetLiver
3094 555773 1972.D24.gz43 247747 F M00054739A:F07 UC2-ColonMetLiver 3095 556019 1981.D01.gz43 247786 F M00054747D:B05 UC2-ColonMetLiver 3096 556019 1981.D01.gz43 247784 F M00054764D:F01 UC2-ColonMetLiver 3097 472196 1981.D03.gz43 247784 F M00054765A:A10 UC2-ColonMetLiver 3098 553158 1981.D03.gz43 247797 F M00054755P:B05 UC2-ColonMetLiver 3100 498662 1981.L03.gz43 247810 F M00054766:B08 UC2-ColonMetLiver 3101 554021 1981.04.gz43 247810 F M00054766:B08 UC2-ColonMetLiver 3102 554021 1981.04.gz43 247810 F M00054760:D07 UC2-ColonMetLiver 3104 448510 1981.P04.gz43 247822 F M00054769:D09 UC2-ColonMetLiver 3104 54890 1981.05.gz43 247832 F M00054765:D09 UC2-ColonMetLiver	3092	560282	1972.C21.gz43_247698		M00054738B:E12	UC2-ColonMetLiver
3095 474309 1972.M24.gz43 247756 F M00054747D:B05 UC2-ColonMetLiver 3096 556019 1981.J01.gz43 247769 F M00054764D:F01 UC2-ColonMetLiver 3097 472196 1981.I02.gz43 247787 F M00054763A:A10 UC2-ColonMetLiver 3098 513632 1981.D03.gz43 247797 F M00054755D:E05 UC2-ColonMetLiver 3099 553158 1981.F03.gz43 247803 F M00054759A:B08 UC2-ColonMetLiver 3100 4986.02 1981.L04.gz43 247810 F M00054766C:B08 UC2-ColonMetLiver 3101 554021 1981.I04.gz43 247810 F M00054766C:B01 UC2-ColonMetLiver 3104 54802 1981.I04.gz43 247822 F M00054766:B02 UC2-ColonMetLiver 3104 54800 1981.I04.gz43 247822 F M00054769E:D12 UC2-ColonMetLiver 3104 54800 1981.I05.gz43 247823 F M00054769E:D05 UC2-ColonMetLiver <td>3093</td> <td>503452</td> <td>1972.L22.gz43_247723</td> <td>F</td> <td>M00054746D:E05</td> <td>UC2-ColonMetLiver</td>	3093	503452	1972.L22.gz43_247723	F	M00054746D:E05	UC2-ColonMetLiver
3096 556019 1981J01.g243 247769 F M00054764D:F01 UC2-ColonMetLiver 3097 472196 1981L02.gz43 247784 F M00054763A:A10 UC2-ColonMetLiver 3098 513632 1981L03.gz43 247795 F M00054755D:E05 UC2-ColonMetLiver 3099 553158 1981F03.gz43 247797 F M00054759A:B08 UC2-ColonMetLiver 3100 488662 1981L03.gz43 247803 F M00054759A:B08 UC2-ColonMetLiver 3101 551693 1981C04.gz43 247816 F M00054769C:B08 UC2-ColonMetLiver 3102 554021 1981.004.gz43 247816 F M00054769B:D12 UC2-ColonMetLiver 3104 448510 1981.004.gz43 247822 F M00054769B:D12 UC2-ColonMetLiver 3105 554294 1981.05.gz43 247832 F M00054769C:D05 UC2-ColonMetLiver 3107 450840 1981.06.gz43 247834 F M00054769C:D05 UC2-ColonMetLiver	3094	555773	1972.D24.gz43_247747	F	M00054739A:F07	UC2-ColonMetLiver
3097 472196 1981.102.gz43_247784 F M00054763A:A10 UC2-ColonMetLiver 3098 513632 1981.D03.gz43_247795 F M00054755D:E05 UC2-ColonMetLiver 3099 553158 1981.F03.gz43_247797 F M00054759A:B08 UC2-ColonMetLiver 3100 498662 1981.L03.gz43_247810 F M0005476C:B08 UC2-ColonMetLiver 3101 551693 1981.C04.gz43_247810 F M0005476C:B08 UC2-ColonMetLiver 3102 554021 1981.004.gz43_247816 F M0005476D:C107 UC2-ColonMetLiver 3103 554042 1981.004.gz43_247822 F M00054769B:D12 UC2-ColonMetLiver 3104 448510 1981.05.gz43_247823 F M00054765D:D09 UC2-ColonMetLiver 3105 554024 1981.K05.gz43_247834 F M00054765D:D05 UC2-ColonMetLiver 3107 450840 1981.106.gz43_247834 F M00054765D:D05 UC2-ColonMetLiver 3107 450840 1981.106.gz43_247834 F M00054765D:D05 U	3095	474309	1972.M24.gz43_247756	F	M00054747D:B05	UC2-ColonMetLiver
3098 513632 1981.D03.gz43 247795 F M00054755D:E05 UC2-ColonMetLiver 3099 553158 1981.F03.gz43 247797 F M00054759A:B08 UC2-ColonMetLiver 3100 498662 1981.L03.gz43 247810 F M00054754D:F11 UC2-ColonMetLiver 3101 551693 1981.C04.gz43 247810 F M00054754D:F11 UC2-ColonMetLiver 3102 554021 1981.I04.gz43 247816 F M00054763C:D07 UC2-ColonMetLiver 3103 554024 1981.P04.gz43 247822 F M00054763E:D12 UC2-ColonMetLiver 3104 448510 1981.P04.gz43 247823 F M00054765D:D05 UC2-ColonMetLiver 3105 554294 1981.R05.gz43 247834 F M00054765D:D05 UC2-ColonMetLiver 3107 450840 1981.R06.gz43 247834 F M00054765:DD05 UC2-ColonMetLiver UC2-ColonMetLiver UC2-ColonMetLiver UC2-ColonMetLiver UC2-ColonMetLiver 3107	3096	556019	1981.J01.gz43_247769	F	M00054764D:F01	UC2-ColonMetLiver
3099 553158 1981.F03.gz43 247797 F M00054759A:B08 UC2-ColonMetLiver 3100 498662 1981.L03.gz43 247803 F M00054766C:B08 UC2-ColonMetLiver 3101 551693 1981.C04.gz43 247810 F M00054754D:F11 UC2-ColonMetLiver 3102 554021 1981.004.gz43 247816 F M00054763C:D07 UC2-ColonMetLiver 3104 448510 1981.P04.gz43 247822 F M00054763D:D12 UC2-ColonMetLiver 3105 558900 1981.105.gz43 247832 F M00054763CF10 UC2-ColonMetLiver 3106 554294 1981.K05.gz43 247834 F M00054763CF10 UC2-ColonMetLiver 3107 450840 1981.R05.gz43 247834 F M00054763CF10 UC2-ColonMetLiver	3097	472196	1981.I02.gz43_247784	F	M00054763A:A10	UC2-ColonMetLiver
3100 498662 1981.1.03 gz43 247803 F M00054766C:B08 UC2-ColonMetLiver	3098	513632	1981.D03.gz43_247795	F	M00054755D:E05	UC2-ColonMetLiver
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3102 554021 1981.104.gz43 247816 F M00054763C:D07 UC2-ColonMetLiver 3103 550402 1981.004.gz43 247822 F M00054769B:D12 UC2-ColonMetLiver 3104 448510 1981.P04.gz43 247823 F M00054769B:D19 UC2-ColonMetLiver 3105 558900 1981.105.gz43 247823 F M00054763C:F10 UC2-ColonMetLiver 3106 554294 1981.K05.gz43 247848 F M00054763C:H04 UC2-ColonMetLiver 3107 450840 1981.106.gz43 247848 F M00054763C:H04 UC2-ColonMetLiver	3100	498662	1981.L03.gz43_247803	F	M00054766C:B08	UC2-ColonMetLiver
3103 550402 1981.004.gz43 247822 F M00054769B:D12 UC2-ColonMetLiver 3104 448510 1981.P04.gz43 247823 F M00054770B:D09 UC2-ColonMetLiver 3105 558900 1981.105.gz43 247832 F M00054763C:F10 UC2-ColonMetLiver 3106 554294 1981.K05.gz43 247834 F M00054763D:D05 UC2-ColonMetLiver 3107 450840 1981.106.gz43 247848 F M00054763C:H04 UC2-ColonMetLiver	3101	551693	1981.C04.gz43_247810	F	M00054754D:F11	UC2-ColonMetLiver
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3106 554294 1981.K05.gz43 247834 F M00054765D:D05 UC2-ColonMetLiver 3107 450840 1981.I06.gz43 247848 F M00054763C:H04 UC2-ColonMetLiver	3104	448510	1981.P04.gz43_247823	F	M00054770B:D09	UC2-ColonMetLiver
3106 554294 1981.K05.gz43 247834 F M00054765D:D05 UC2-ColonMetLiver 3107 450840 1981.I06.gz43 247848 F M00054763C:H04 UC2-ColonMetLiver	3105	558900	1981.I05.gz43_247832	F	M00054763C:F10	UC2-ColonMetLiver
	3106	554294	1981.K05.gz43_247834	F	M00054765D:D05	
3108 561487 1981.J07.gz43 247865 F M00054765B:C03 UC2-ColonMetLiver	3107	450840	1981.I06.gz43_247848	F	M00054763C:H04	UC2-ColonMetLiver
The state of the s	3108	561487	1981.J07.gz43_247865	F	M00054765B:C03	UC2-ColonMetLiver

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SEQ					
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NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
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3110	528775	1981.P07.gz43_247871	F	M00054770C:A04	UC2-ColonMetLiver
3111	553918	1981.B08.gz43_247873	F	M00054753C:H02	UC2-ColonMetLiver
3112	477046	1981.B09.gz43_247889	F	M00054753D:A03	UC2-ColonMetLiver
3113	551289	1981.D09.gz43_247891	F	M00054756A:C12	UC2-ColonMetLiver
3114	263800	1981.G09.gz43_247894	F	M00054760D:B03	UC2-ColonMetLiver
3115	495942	1981.A10.gz43_247904	F	M00054752B:H06	UC2-ColonMetLiver
3116	556183	1981.H10.gz43_247911	F	M00054762A:D09	UC2-ColonMetLiver
3117	448357	1981.N10.gz43_247917	F	M00054768D:A01	UC2-ColonMetLiver
3118	554161	1981.P10.gz43_247919	F	M00054770C:C04	UC2-ColonMetLiver
3119	256179	1981.B11.gz43_247921	F	M00054753D:C12	UC2-ColonMetLiver
3120	555478	1981.P11.gz43_247935	F	M00054770C:D05	UC2-ColonMetLiver
3121	554581	1981.P12.gz43_247951	F	M00054770C:F10	UC2-ColonMetLiver
3122	553922	1981.B13.gz43_247953	F	M00054753D:H10	UC2-ColonMetLiver
3123	554336	1981.E13.gz43_247956	F	M00054758A:F03	UC2-ColonMetLiver
3124	560529	1981.F13.gz43_247957	F	M00054759C:G10	UC2-ColonMetLiver
3125	558900	1981.H14.gz43_247975	F	M00054762B:F02	UC2-ColonMetLiver
3126	447667	1981.M14.gz43_247980	F	M00054767C:D03	UC2-ColonMetLiver
3127	551288	1981.D15.gz43 247987	F	M00054756C:C08	UC2-ColonMetLiver
3128	122169	1981.E15.gz43_247988	F	M00054758B:C10	UC2-ColonMetLiver
3129	551617	1981.H15.gz43 247991	F	M00054762B:F07	UC2-ColonMetLiver
3130	260558	1981.L15.gz43_247995	F	M00054766D:H02	UC2-ColonMetLiver
3131	558609	1981.B16.gz43 248001	F	M00054754B:F04	UC2-ColonMetLiver
3132	554352	1981.E16.gz43 248004	F	M00054758B:D03	UC2-ColonMetLiver
3133	556064	1981.F17.gz43 248021	F	M00054759D:E01	UC2-ColonMetLiver
3134	508126	1981.K17.gz43 248026	F	M00054766A:H10	UC2-ColonMetLiver
3135	549576	1981.M17.gz43 248028	F	M00054767C:H06	UC2-ColonMetLiver
3136	554000	1981.A18.gz43 248032	F	M00054753A:A05	UC2-ColonMetLiver
3137	554049	1981.E18.gz43 248036	F	M00054758B:H03	UC2-ColonMetLiver
3138	558503	1981.H18.gz43 248039	F	M00054762C:A12	UC2-ColonMetLiver
3139	451993	1981.D19.gz43 248051	F	M00054756D:F05	UC2-ColonMetLiver
3140	558949	1981.O19.gz43 248062	F	M00054770A:C06	UC2-ColonMetLiver
3141	553131	1981.J20.gz43 248073	F	M00054765C:F10	UC2-ColonMetLiver
3142	555343	1981.D21.gz43 248083	F	M00054757A:H07	UC2-ColonMetLiver
3143	512432	1981.N21.gz43 248093	F	M00054769A:G10	UC2-ColonMetLiver
3144	550580	1981.B22.gz43 248097	F	M00054754C:H09	UC2-ColonMetLiver
3145	551117	1981.F22.gz43 248101	F	M00054760A:A12	UC2-ColonMetLiver
3146	556475	1981.L22.gz43 248107	F	M00054767A:F08	UC2-ColonMetLiver
3147	552549	1981.M22.gz43_248108	F	M00054767D:G09	UC2-ColonMetLiver
3148	470351	1981.P23.gz43_248127	F	M00054771A:E01	UC2-ColonMetLiver
3149	561535	1981.F24.gz43_248133	F	M00054760A:D10	UC2-ColonMetLiver
3150	1093	1981.G24.gz43_248134	F	M00054761C:E02	UC2-ColonMetLiver

Table 2

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3152	555958	1982.F01.gz43_248149	F	M00054778A:D01	UC2-ColonMetLiver
3153	4927 7 9	1982.A02.gz43_248160	F	M00054771B;F12	UC2-ColonMetLiver
3154	553848	1982.P02.gz43_248175	F	M00054806A:G04	UC2-ColonMetLiver
3155	555686	1982.C03.gz43_248178	F	M00054774B:A07	UC2-ColonMetLiver
3156	555524	1982.H03.gz43_248183	F	M00054780B:B06	UC2-ColonMetLiver
3157	551068	1982.A04.gz43_248192	F	M00054771C:A11	UC2-ColonMetLiver
3158	552183	1982.O04.gz43_248206	F	M00054805A:H04	UC2-ColonMetLiver
3159	551003	1982.F05.gz43_248213	F	M00054778A:F08	UC2-ColonMetLiver
3160	465207	1982.G05.gz43_248214	F	M00054779B;A07	UC2-ColonMetLiver
3161	533991	1982.H05.gz43_248215	F	M00054780B:E03	UC2-ColonMetLiver
3162	553868	1982.K05.gz43_248218	F	M00054786D:F08	UC2-ColonMetLiver
3163	553774	1982.B06.gz43_248225	F	M00054772D:G12	UC2-ColonMetLiver
3164	554072	1982.C06.gz43_248226	F	M00054774B:G08	UC2-ColonMetLiver
3165	554828	1982.E06.gz43_248228	F	M00054776C:G06	UC2-ColonMetLiver
3166	559872	1982.G06.gz43_248230	F	M00054779B:B07	UC2-ColonMetLiver
3167	491570	1982.H06.gz43_248231	F	M00054780B:G10	UC2-ColonMetLiver
3168	275	1982.M06.gz43_248236	F	M00054802C:A07	UC2-ColonMetLiver
3169	552019	1982.C07.gz43_248242	F	M00054774C:A03	UC2-ColonMetLiver
3170	556183	1982.G07.gz43_248246	F	M00054779B:B11	UC2-ColonMetLiver
3171	552535	1982.J07.gz43_248249	F	M00054784C:D11	UC2-ColonMetLiver
3172	551718	1982.A08.gz43_248256	F	M00054771D:G01	UC2-ColonMetLiver
3173	551169	1982.B08.gz43_248257	F	M00054773A:E09	UC2-ColonMetLiver
3174	551080	1982.F08.gz43_248261	F	M00054778B:A02	UC2-ColonMetLiver
3175	517146	1982.P08.gz43_248271	F	M00054806C:C12	UC2-ColonMetLiver
3176	402147	1982.B09.gz43_248273	F	M00054773A:H05	UC2-ColonMetLiver
3177	557269	1982.D09.gz43_248275	F	M00054776A:D10	UC2-ColonMetLiver
3178	539353	1982.N09.gz43_248285	F	M00054804A:H04	UC2-ColonMetLiver
3179	560520	1982.D10.gz43_248291	F	M00054776A:F01	UC2-ColonMetLiver
3180	575924	1982.I10.gz43_248296	F	M00054782D:D12	UC2-ColonMetLiver
3181	554582	1982.J10.gz43_248297	F	M00054784D:B01	UC2-ColonMetLiver
3182	555696	1982.M10.gz43_248300	F	M00054802D:A09	UC2-ColonMetLiver
3183	556856	1982.O10.gz43_248302	F	M00054805B:E06	UC2-ColonMetLiver
3184	447926	1982.D11.gz43_248307	F	M00054776A:F07	UC2-ColonMetLiver
3185	553743	1982.E11.gz43_248308	F	M00054776D:G09	UC2-ColonMetLiver
3186	553979	1982.N11.gz43_248317	F	M00054804B:E07	UC2-ColonMetLiver
3187	497912	1982.O11.gz43_248318	F	M00054805B:E11	UC2-ColonMetLiver
3188	412621	1982.D12.gz43_248323	F	M00054776A:G01	UC2-ColonMetLiver
3189	497477	1982.M12.gz43_248332	F	M00054802D:C02	UC2-ColonMetLiver
3190	552188	1982.J13.gz43_248345	F	M00054785B:B07	UC2-ColonMetLiver
3191	456052	1982.M13.gz43_248348	F	M00054802D:C03	UC2-ColonMetLiver
3192	143346	1982.H14.gz43_248359	F	M00054780C:G11	UC2-ColonMetLiver

Table 2

	able 2				
SEQ					
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NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
3193	553237	1982.N14.gz43_248365	F	M00054804C:F04	UC2-ColonMetLiver
3194	556856	1982.O14.gz43_248366	F	M00054805B:G02	UC2-ColonMetLiver
3195	586600	1982.A15.gz43_248368	F	M00054772B:B01	UC2-ColonMetLiver
3196	562263	1982.J15.gz43_248377	F	M00054785C:C02	UC2-ColonMctLiver
3197	555462	1982.C16.gz43_248386	F	M00054775A:D07	UC2-ColonMctLiver
3198	394168	1982.O16.gz43_248398	F	M00054805B:G10	UC2-ColonMetLiver
3199	472672	1982.A17.gz43_248400	F	M00054772B:D04	UC2-ColonMetLiver
3200	595181	1982.C17.gz43_248402	F	M00054775A:G03	UC2-ColonMetLiver
3201	554627	1982.G17.gz43_248406	F	M00054779D:F07	UC2-ColonMctLiver
3202	552325	1982.I17.gz43_248408	F	M00054783C:C03	UC2-ColonMctLiver
3203	462511	1982.J17.gz43_248409	F	M00054785C:G11	UC2-ColonMctLiver
3204	552813	1982.K17.gz43_248410	F	M00054788C:G04	UC2-ColonMetLiver
3205	175758	1982.M17.gz43_248412	F	M00054803A:D08	UC2-ColonMctLiver
3206	493085	1982.A18.gz43_248416	F	M00054772B:F03	UC2-ColonMctLiver
3207	557834	1982.B18.gz43 248417	F	M00054773C:F09	UC2-ColonMetLiver
3208	548998	1982.F18.gz43 248421	F	M00054778C:F09	UC2-ColonMetLiver
3209	551425	1982.G18.gz43 248422	F	M00054779D:F08	UC2-ColonMetLiver
3210	490414	1982.K18.gz43 248426	F	M00054789A:A02	UC2-ColonMetLiver
3211	551553	1982.M18.gz43 248428	F	M00054803A:E10	UC2-ColonMetLiver
3212	555571	1982.N18.gz43 248429	F	M00054804D:D07	UC2-ColonMetLiver
3213	448629	1982.B19.gz43_248433	F	M00054773C:F10	UC2-ColonMetLiver
3214	552005	1982.J19.gz43_248441	F	M00054785D:A07	UC2-ColonMetLiver
3215	554116	1982.M20.gz43 248460	F	M00054803B:B12	UC2-ColonMetLiver
3216	505792	1982.H21.gz43_248471	F	M00054781A:H09	UC2-ColonMctLiver
3217	561422	1982.J21.gz43_248473	F	M00054786A:G11	UC2-ColonMetLiver
3218	551755	1982.M24.gz43_248524	F	M00054803C:G01	UC2-ColonMetLiver
3219	550063	1945.J03.gz43_248963	F	M00054513A:B08	UC2-ColonMetLiver
3220	388055	1945.D04.gz43_248973	F	M00054506B:A07	UC2-ColonMetLiver
3221	572807	1945.C05.gz43_248988	F	M00054505A:G12	UC2-ColonMetLiver
3222	455814	1945.J05.gz43_248995	F	M00054513A:F09	UC2-ColonMetLiver
3223	452471	1945.H07.gz43_249025	F	M00054510C:G07	UC2-ColonMetLiver
3224	551518	1945.J07.gz43_249027	F	M00054513B:E11	UC2-ColonMetLiver
3225	555660	1945.J09.gz43_249059	F	M00054513C:A01	UC2-ColonMetLiver
3226	524721	1945.B11.gz43_249083	F	M00054504B:A04	UC2-ColonMetLiver
3227	550986	1945.D11.gz43_249085	F	M00054506B:H01	UC2-ColonMetLiver
3228	499696	1945.G11.gz43_249088	F	M00054509C:G01	UC2-ColonMetLiver
3229	554611	1945.N11.gz43_249095	F	M00054519A;C04	UC2-ColonMetLiver
3230	557681	1945.F12.gz43_249103	F	M00054508B:E08	UC2-ColonMetLiver
3231	550678	1945.J13.gz43_249123	F	M00054513D:F04	UC2-ColonMetLiver
3232	554246	1945.K13.gz43_249124	F	M00054515B:E11	UC2-ColonMetLiver
3233	550129	1945.F15.gz43_249151	F	M00054508C:B04	UC2-ColonMetLiver
3234	555771	1945.F17.gz43 249183	F	M00054508C:C08	UC2-ColonMetLiver

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Table 2

	able 2				
SEQ					
ID			ORIE		
NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
3235	572992	1945.O18.gz43_249208	F	M00054521D:H01	UC2-ColonMetLiver
3236	461313	1945.J21.gz43_249251	F	M00054514C:B01	UC2-ColonMetLiver
3237	554456	1945.L21.gz43_249253	F	M00054516D:F09	. UC2-ColonMetLiver
3238	547866	1945.F23.gz43_249279	F	M00054509A:C01	UC2-ColonMetLiver
3239	458974	1945.P23.gz43_249289	F	M00054523C:A11	UC2-ColonMetLiver
3240	554839	1945.C24.gz43_249292	F	M00054506A:D05	UC2-ColonMetLiver
3241	487658	1945.I24.gz43_249298	F	M00054512D:H08	UC2-ColonMetLiver
3242	556940	1946.H01.gz43_249313	F	M00054533C:B04	UC2-ColonMetLiver
3243	493575	1946.N03.gz43_249351	F	M00054541C:C10	UC2-ColonMetLiver
3244	466020	1946.H05.gz43_249377	F	M00054533C:E07	UC2-ColonMetLiver
3245	552006	1946.C06.gz43_249388	F	M00054527B:A07	UC2-ColonMetLiver
3246	551157	1946.E06.gz43_249390	F	M00054529C:D11	UC2-ColonMetLiver
3247	559574	1946.G06.gz43_249392	F	M00054532D:A01	UC2-ColonMetLiver
3248	556216	1946.I06.gz43_249394	F	M00054534D:D02	UC2-ColonMetLiver
3249	559676	1946.B07.gz43_249403	F	M00054525D:H05	UC2-ColonMetLiver
3250	550018	1946.J07.gz43_249411	F	M00054536B:A03	UC2-ColonMetLiver
3251	526984	1946.G08.gz43_249424	F	M00054532D:E07	UC2-ColonMetLiver
3252	481594	1946.N09.gz43_249447	F	M00054541C:F11	UC2-ColonMetLiver
3253	550618	1946.P09.gz43_249449	F	M00054544B:E03	UC2-ColonMetLiver
3254	454336	1946.A10.gz43_249450	F	M00054523D:G09	UC2-ColonMetLiver
3255	550370	1946.J10.gz43_249459	F	M00054536B:D02	UC2-ColonMetLiver
3256	464154	1946.M10.gz43_249462	F	M00054540A:H07	UC2-ColonMetLiver
3257	550475	1946.B11.gz43_249467	F	M00054526A:E04	UC2-ColonMetLiver
3258	584071	1946.F11.gz43_249471	F	M00054531D:F05	UC2-ColonMetLiver
3259	550730	1946.J11.gz43_249475	F	M00054536B:F08	UC2-ColonMetLiver
3260	552638	1946.O12.gz43_249496	F	M00054543B;E06	UC2-ColonMetLiver
3261	560014	1946.L13.gz43_249509	F	M00054538D:H11	UC2-ColonMetLiver
3262	553338	1946.I14.gz43_249522	F	M00054535A:G03	UC2-ColonMetLiver
3263	556809	1946.B15.gz43_249531	F	M00054526B:C09	UC2-ColonMetLiver
3264	498433	1946.H15.gz43_249537	F	M00054534A:B06	UC2-ColonMetLiver
3265	455821	1946.K16.gz43_249556	F	M00054538A:E10	UC2-ColonMetLiver
3266	82864	1946.J17.gz43_249571	F	M00054536C:D12	UC2-ColonMetLiver
3267	560519	1946.L17.gz43 249573	F	M00054539B;D06	UC2-ColonMetLiver
3268	409262	1946.N17.gz43_249575	F	M00054542B:B01	UC2-ColonMetLiver
3269	560717	1946.P17.gz43_249577	F	M00054544C:F04	UC2-ColonMetLiver
3270	550172	1946.K18.gz43_249588	F	M00054538B:B05	UC2-ColonMetLiver
3271	552753	1946.O18.gz43_249592	F	M00054543C:F01	UC2-ColonMetLiver
3272	549734	1946.L19.gz43_249605	F	M00054539B:G03	UC2-ColonMetLiver
3273	550714	1946.G20.gz43_249616	F	M00054533B:F04	UC2-ColonMetLiver
3274	585380	1946.D21.gz43_249629	F	M00054529A:H01	UC2-ColonMetLiver
3275	550204	1946.F21.gz43_249631	F	M00054532B:C05	UC2-ColonMetLiver
3276	560205	1946.I22.gz43_249650	F	M00054535B:H08	UC2-ColonMetLiver

Table 2

_ T	able 2				
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NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
3277	312036	1946.E23.gz43_249662	F	M00054530D:C10	UC2-ColonMetLiver
3278	553002	1946.E24.gz43_249678	F	M00054530D:H07	UC2-ColonMetLiver
3279	394567	1948.001.gz43_249704	F	M00054583D:E04	UC2-ColonMetLiver
3280	468613	1948.C02.gz43_249708	F	M00054569D:E01	UC2-ColonMetLiver
3281	378610	1948.I02.gz43_249714	F	M00054576D:C07	UC2-ColonMetLiver
3282	559752	1948.M02.gz43_249718	F	M00054581B:A01	UC2-ColonMetLiver
3283	550804	1948.K03.gz43_249732	F	M00054579A:G10	UC2-ColonMetLiver
3284	549956	1948.O03.gz43_249736	F	M00054584A:A07	UC2-ColonMetLiver
3285	556065	1948.A04.gz43_249738	F	M00054567C:B03	UC2-ColonMetLiver
3286	550397	1948.M04.gz43_249750	F	M00054581B:D03	UC2-ColonMetLiver
3287	550874	1948.M05.gz43_249766	F	M00054581B:G10	UC2-ColonMetLiver
3288	533812	1948.O05.gz43_249768	F	M00054584A:B03	UC2-ColonMetLiver
3289	549597	1948.C08.gz43_249804	F	M00054570A:F02	UC2-ColonMetLiver
3290	553749	1948.K08.gz43_249812	F	M00054579B:D10	UC2-ColonMetLiver
3291	446814	1948.G10.gz43_249840	F	M00054574D:F11	UC2-ColonMetLiver
3292	555000	1948.M10.gz43_249846	F	M00054581D:C12	UC2-ColonMetLiver
3293	554887	1948.O10.gz43_249848	F	M00054584B:A03	UC2-ColonMetLiver
3294	561876	1948.E11.gz43_249854	F	M00054572C:E09	UC2-ColonMetLiver
3295	556488	1948.M11.gz43_249862	F	M00054581D:D01	UC2-ColonMetLiver
3296	560080	1948.G12.gz43_249872	· F	M00054575A:B06	UC2-ColonMetLiver
3297	556336	1948.M12.gz43_249878	F	M00054581D:E04	UC2-ColonMetLiver
3298	558729	1948.I13.gz43_249890	F	M00054577B:A09	UC2-ColonMetLiver
3299	549716	1948.A14.gz43_249898	F	M00054568A:G02	UC2-ColonMetLiver
3300	568467	1948.M14.gz43_249910	F	M00054582A:A05	UC2-ColonMetLiver
3301	550694	1948.N14.gz43_249911	F	M00054583A:F05	UC2-ColonMetLiver
3302	549994	1948.M15.gz43_249926	F	M00054582A:A07	UC2-ColonMetLiver
3303	549858	1948.A16.gz43_249930	F	M00054568A:H03	UC2-ColonMetLiver
3304	554793	1948.A17.gz43_249946	F	M00054568B:A07	UC2-ColonMetLiver
3305	466020	1948.O17.gz43_249960	F	M00054584D:C01	UC2-ColonMetLiver
3306	509202	1948.I19.gz43_249986	F	M00054577B:F01	UC2-ColonMetLiver
3307	450963	1948.A21.gz43_250010	F	M00054568C:D06	UC2-ColonMetLiver
3308	451049	1948.C21.gz43_250012	F	M00054571A:B10	UC2-ColonMetLiver
3309	528404	1948.K21.gz43_250020	F	M00054580A:B11	UC2-ColonMetLiver
3310	160320	1948.O21.gz43_250024	F	M00054585A:E07	UC2-ColonMetLiver
3311	492139	1948.K22.gz43_250036	F	M00054580A:C10	UC2-ColonMetLiver
3312	557656	1948.A23.gz43_250042	F	M00054568C:G12	UC2-ColonMetLiver
3313	570248	1948.K24.gz43_250068	F	M00054580A:D04	UC2-ColonMetLiver
3314	561836	1957.G01.gz43_250080	F	M00054594A:D04	UC2-ColonMetLiver
3315	558332	1957.J01.gz43_250083	F	M00054597A:C11	UC2-ColonMetLiver
3316	558981	1957.J02.gz43_250099	F	M00054597A:D08	UC2-ColonMetLiver
3317	535129	1957.B03.gz43_250107	F	M00054587C:G02	UC2-ColonMetLiver
3318	549781	1957.J03.gz43_250115	F	M00054597A:G08	UC2-ColonMetLiver

Table 2

T	able 2				
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NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
3319	555920	1957.C04.gz43_250124	F	M00054589A:D09	UC2-ColonMetLiver
3320	550044	1957.E04.gz43_250126	F	M00054591C:A01	UC2-ColonMetLiver
3321	447356	1957.I04.gz43_250130	F	M00054596B:C01	UC2-ColonMetLiver
3322	560751	1957.A05.gz43_250138	F	M00054586C:H02	UC2-ColonMetLiver
3323	550322	1957.B05.gz43_250139	F	M00054587D:D04	UC2-ColonMetLiver
3324	416884	1957.N05.gz43_250151	F	M00054602A:E06	UC2-ColonMetLiver
3325	559857	1957.A06.gz43_250154	F	M00054586D:A03	UC2-ColonMetLiver
3326	446164	1957.C06.gz43_250156	F	M00054589B;A07	UC2-ColonMctLiver
3327	550088	1957.D06.gz43_250157	F	M00054590B;B11	UC2-ColonMetLiver
3328	554764	1957.E06.gz43_250158	F	M00054591C:H09	UC2-ColonMetLiver
3329	552907	1957.007.gz43_250184	F	M00054603A:G06	UC2-ColonMetLiver
3330	498827	1957.D08.gz43_250189	F	M00054590C:A03	UC2-ColonMetLiver
3331	498777	1957.G08.gz43_250192	F	M00054594C:E06	UC2-ColonMetLiver
3332	559883	1957.L08.gz43_250197	F	M00054599D:B07	UC2-ColonMetLiver
3333	554869	1957.M08.gz43_250198	F	M00054600D:H07	UC2-ColonMetLiver
3334	559419	1957.A09.gz43_250202	F	M00054586D:G01	UC2-ColonMetLiver
3335	473859	1957.F09.gz43_250207	F	M00054593B:B03	UC2-ColonMetLiver
3336	555949	1957.M09.gz43_250214	F	M00054601A:E08	UC2-ColonMetLiver
3337	552913	1957.O09.gz43_250216	F	M00054603A:G12	UC2-ColonMetLiver
3338	551463	1957.C10.gz43_250220	F	M00054589B:F05	UC2-ColonMetLiver
3339	550855	1957.B11.gz43_250235	F	M00054588A:G08	UC2-ColonMetLiver
3340	526733	1957.O11.gz43_250248	F	M00054603B:B04	UC2-ColonMetLiver
3341	557970	1957.F12.gz43_250255	F	M00054593B:E03	UC2-ColonMetLiver
3342	549665	1957.H12.gz43_250257	F	M00054595C:G06	UC2-ColonMetLiver
3343	446503	1957.M12.gz43_250262	F	M00054601B:D08	UC2-ColonMetLiver
3344	549599	1957.P12.gz43_250265	F	M00054604C:F03	UC2-ColonMetLiver
3345	558413	1957.A13.gz43_250266	F	M00054587A:A08	UC2-ColonMetLiver
3346	553331	1957.B13.gz43_250267	F	M00054588B;B12	UC2-ColonMetLiver
3347	549609	1957.I14.gz43_250290	F	M00054596C:F09	UC2-ColonMetLiver
3348	455298	1957.I15.gz43_250306	F	M00054596C:F10	UC2-ColonMetLiver
3349	563514	1957.P15.gz43_250313	F	M00054604D:A04	UC2-ColonMetLiver
3350	471181	1957.A16.gz43_250314	F	M00054587A:F06	UC2-ColonMetLiver
3351	487176	1957.N16.gz43_250327	F	M00054602C:C12	UC2-ColonMetLiver
3352	561130	1957.I17.gz43_250338	F	M00054596C:G09	UC2-ColonMetLiver
3353	461718	1957.J17.gz43_250339	F	M00054597D:E12	UC2-ColonMetLiver
3354	559554	1957.L17.gz43_250341	F	M00054600A:G12	UC2-ColonMetLiver
3355	452506	1957.A18.gz43_250346	F	M00054587A:F09	UC2-ColonMetLiver
3356	551975	1957.D18.gz43_250349	F	M00054590D:C12	UC2-ColonMetLiver
3357	485029	1957.H18.gz43_250353	F	M00054595D:A10	UC2-ColonMetLiver
3358	549816	1957.L18.gz43_250357	F	M00054600A:H03	UC2-ColonMetLiver
3359	408130	1957.O18.gz43_250360	F	M00054603C:G09	UC2-ColonMetLiver
3360	558927	1957.P18.gz43_250361	F	M00054605A:A03	UC2-ColonMetLiver

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3389 552823 1958.B07.gz43_250555 F M00054607A:G02 UC2-ColonMetLi	/er
3390 395411 1958.M07.gz43_250566 F M00054620D:G11 UC2-ColonMetLi	/er
3391 10397 1958.N07.gz43_250567 F M00054622A:D06 UC2-ColonMetLi	er
3392 551250 1958.P07.gz43_250569 F M00054625B:B02 UC2-ColonMetLi	/er
3393 561892 1958.C08.gz43_250572 F M00054609A:F01 UC2-ColonMetLi	/er
3394 549347 1958.F08.gz43 250575 F M00054612A:D12 UC2-ColonMetLi	/er
3395 561068 1958.N08.gz43_250583 F M00054622A;H01 UC2-ColonMetLi	/er
3396 549912 1958.C09.gz43_250588 F M00054609A:H04 UC2-ColonMetLi	/er
3397 549114 1958.F10.gz43_250607 F M00054612B:B11 UC2-ColonMetLi	/ег
3398 548965 1958.J10.gz43_250611 F M00054617B:A09 UC2-ColonMetLi	/er
3399 491644 1958.N10.gz43_250615 F M00054622B:F05 UC2-ColonMetLi	/er
3400 550018 1958.G11.gz43_250624 F M00054613B:H04 UC2-ColonMetLi	/er
3401 556530 1958.C12.gz43 250636 F M00054609B;E01 UC2-ColonMetLi	/er
3402 562749 1958.L12.gz43_250645 F M00054619D:C10 UC2-ColonMetLi	

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Table 2

T	able 2				
SEQ					
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NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
3403	556308	1958.N12.gz43_250647	F	M00054622B:H09	UC2-ColonMetLiver
3404	551305	1958.O12.gz43_250648	F	M00054623D:C12	UC2-ColonMctLiver
3405	507349	1958.C13.gz43_250652	F	M00054609B:H11	UC2-ColonMetLiver
3406	549626	1958.E13.gz43_250654	F	M00054611B:F12	UC2-ColonMetLiver
3407	549388	1958.J13.gz43_250659	F	M00054617B:D06	UC2-ColonMetLiver
3408	478511	1958.E14.gz43_250670	F	M00054611B:G09	UC2-ColonMctLiver
3409	555371	1958.K14.gz43_250676	F	M00054618C:H02	UC2-ColonMetLiver
3410	553758	1958.I15.gz43_250690	F	M00054616A:H01	UC2-ColonMetLiver
3411	553204	1958.L15.gz43_250693	F	M00054620A:C09	UC2-ColonMetLiver
3412	497477	1958.H16.gz43_250705	F	M00054615B:E03	UC2-ColonMetLiver
3413	481360	1958.I16.gz43 250706	F	M00054616A:H03	UC2-ColonMetLiver
3414	552437	1958.K16.gz43 250708	F	M00054618D:D04	UC2-ColonMetLiver
3415	550164	1958.O16.gz43 250712	F	M00054624A:B11	UC2-ColonMetLiver
3416	555660	1958.E18.gz43 250734	F	M00054611C:E01	UC2-ColonMetLiver
3417		1958.N18.gz43 250743	F	M00054622D:C02	UC2-ColonMetLiver
3418		1958.B19.gz43 250747	F	M00054608B:D08	UC2-ColonMetLiver
3419		1958.K19.gz43 250756	F	M00054618D:E06	UC2-ColonMetLiver
3420	451009	1958.K20,gz43 250772	F	M00054618D:E11	UC2-ColonMetLiver
3421	552194	1958.L20.gz43 250773	F	M00054620B:B10	UC2-ColonMetLiver
3422	451025	1958,M20,gz43 250774	F	M00054621C:G03	UC2-ColonMetLiver
3423	551463	1958.N20.gz43 250775	F	M00054622D:D10	UC2-ColonMetLiver
3424	553028	1958.G21.gz43_250784	F	M00054613D:H09	UC2-ColonMetLiver
3425	485504	1958.N21.gz43 250791	F	M00054622D:F05	UC2-ColonMetLiver
3426	562712	1958.P21.gz43_250793	F	M00054625D:F06	UC2-ColonMetLiver
3427	490152	1958.C22.gz43 250796	F	M00054609D:E12	UC2-ColonMetLiver
3428	552364	1958.F22.gz43 250799	F	M00054612D;B12	UC2-ColonMetLiver
3429	448276	1958.L22.gz43 250805	F	M00054620C;C08	UC2-ColonMetLiver
3430	556835	1958.N22.gz43 250807	F	M00054622D;G11	UC2-ColonMetLiver
3431	461734	1958.P22.gz43 250809	F	M00054625D:H07	UC2-ColonMetLiver
3432	557947	1958.H23.gz43_250817	F	M00054615C:D09	UC2-ColonMetLiver
3433	560491	1958.L23.gz43_250821	F	M00054620C;C12	UC2-ColonMetLiver
3434	554810	1958.O23.gz43_250824	F	M00054624D;B06	UC2-ColonMetLiver
3435	549829	1958,C24,gz43 250828	F	M00054609D:H06	UC2-ColonMctLiver
3436	518049	1958.M24.gz43 250838	F	M00054621D:A10	UC2-ColonMetLiver
3437	484669	1777.O01.gz43_251861	F	M00042831D:G06	UC2-PrimColon
3438	537506	1777.C03.gz43 251881	F	M00042831D:000	UC2-PrimColon
3439	545162	1777.F03.gz43_251884	F	M00042811C:B00	UC2-PrimColon
3440	456793	1777.C11.gz43 252009	F	M00042810B:C08	UC2-PrimColon
3441	448340	1777.E11.gz43 252011	F	M00042812B:C01	UC2-PrimColon
3442	448793	1777.E13.gz43_252043	F	M00042815C:C02	UC2-PrimColon
3443	449718	1777.H14.gz43 252062	F	M00042819C:H02	UC2-PrimColon
3444			F		
13444	735534	1777.E17.gz43 252107	r	M00042816A:C09	UC2-PrimColon

Table 2

1:	able 2				
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3445	449258	1777.K17.gz43_252113	F	M00042825A:B05	UC2-PrimColon
3446	485020	1777.018.gz43_252133	F	M00042833C:G05	UC2-PrimColon
3447	649735	1777.A23.gz43_252199	F	M00042809D:C12	UC2-PrimColon
3448	498504	1923.B01.gz43_252616	F	M00042352D:F11	UC2-ColonMetLiver
3449	424723	1923.C01.gz43_252617	F	M00042355A:H09	UC2-ColonMetLiver
3450	451124	1923.E01.gz43_252619	F	M00042444C:E02	UC2-ColonMetLiver
3451	553312	1923.O01.gz43_252629	F	M00042516D:H08	UC2-ColonMetLiver
3452	257547	1923.D02.gz43_252634	F	M00042442B:A07	UC2-ColonMetLiver
3453	492292	1923.J02.gz43_252640	F	M00042455B:G09	UC2-ColonMetLiver
3454	638444	1923.002.gz43_252645	F	M00042516D:H09	UC2-ColonMetLiver
3455	451009	1923.C03.gz43 252649	F	M00042355B:B07	UC2-ColonMetLiver
3456	451518	1923.I03.gz43_252655	F	M00042453C:D12	UC2-ColonMetLiver
3457	450255	1923.J03.gz43_252656	F	M00042455C:D11	UC2-ColonMetLiver
3458	556804	1923.M03.gz43_252659	F	M00042460B:D10	UC2-ColonMetLiver
3459	497400	1923.C04.gz43_252665	F	M00042355B:E10	UC2-ColonMetLiver
3460	451618	1923.H04.gz43_252670	F	M00042451D:H10	UC2-ColonMetLiver
3461	490890	1923.J04.gz43_252672	F	M00042455C:E04	UC2-ColonMetLiver
3462	492982	1923.P04.gz43_252678	F	M00042519D:H07	UC2-ColonMetLiver
3463	498777	1923.A05.gz43_252679	F	M00042351C:G01	UC2-ColonMetLiver
3464	493575	1923.B05.gz43_252680	F	M00042353B:A11	UC2-ColonMctLiver
3465	556916	1923.L05.gz43_252690	F	M00042459B:F03	UC2-ColonMctLiver
3466	551755	1923.N05.gz43_252692	F	M00042460D:H06	UC2-ColonMetLiver
3467	495074	1923.O05.gz43_252693	F	M00042517C:B04	UC2-ColonMetLiver
3468	487745	1923.E06.gz43_252699	F	M00042445A:B04	UC2-ColonMetLiver
3469	448663	1923.F06.gz43_252700	F	M00042447C;H10	UC2-ColonMetLiver
3470	468729	1923.L06.gz43_252706	F	M00042459B:F12	UC2-ColonMetLiver
3471	451470	1923.C07.gz43_252713	F	M00042355C:G09	UC2-ColonMetLiver
3472	489426	1923.K07.gz43_252721	F	M00042457D:D02	UC2-ColonMetLiver
3473	450302	1923.F08.gz43_252732	F	M00042447D:E04	UC2-ColonMetLiver
3474	559575	1923.M08.gz43_252739	F	M00042460B:G12	UC2-ColonMetLiver
3475	496446	1923.A09.gz43_252743	F	M00042351D:D02	UC2-ColonMetLiver
3476	492110	1923.F09.gz43_252748	F	M00042447D:G10	UC2-ColonMetLiver
3477	495941	1923.G09.gz43_252749	F	M00042450A:D02	UC2-ColonMetLiver
3478	446621	1923.J09.gz43_252752	F	M00042455D:H08	UC2-ColonMetLiver
3479	562115	1923.N09.gz43_252756	F	M00042516A:A10	UC2-ColonMetLiver
3480	494300	1923.009.gz43_252757	F	M00042517D:A12	UC2-ColonMetLiver
3481	451429	1923.A10.gz43_252759	F	M00042351D:F08	UC2-ColonMetLiver
3482	486912	1923.H10.gz43_252766	F	M00042452C:A09	UC2-ColonMetLiver
3483	456492	1923.J10.gz43_252768	F	M00042456A:C08	UC2-ColonMetLiver
3484	562060	1923.N10.gz43_252772	F	M00042516A:C08	UC2-ColonMetLiver
3485	4244	1923.P10.gz43_252774	F	M00042520A:F09	UC2-ColonMetLiver
3486	451429	1923.A11.gz43_252775	F	M00042351D:H05	UC2-ColonMetLiver

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	able 2				
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NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
3487	449974	1923.H11.gz43_252782	F	M00042452C:C10	UC2-ColonMetLiver
3488	59512	1923.K11.gz43_252785	F	M00042458B;B08	UC2-ColonMetLiver
3489	550362	1923.M11.gz43_252787	F	M00042460C:A02	UC2-ColonMetLiver
3490	211273	1923.O11.gz43_252789	F	M00042517D:G04	UC2-ColonMetLiver
3491	450400	1923.I12.gz43_252799	F	M00042454A:F02	UC2-ColonMetLiver
3492	562627	1923.L12.gz43_252802	F	M00042459C:E06	UC2-ColonMetLiver
3493	494393	1923.B13.gz43_252808	F	M00042353D:B08	UC2-ColonMetLiver
3494	451049	1923.F13.gz43_252812	F	M00042448B:C04	UC2-ColonMetLiver
3495	558729	1923.K13.gz43_252817	F	M00042458B:G05	UC2-ColonMetLiver
3496	526733	1923.L13.gz43_252818	F	M00042459C:G02	UC2-ColonMctLiver
3497	558559	1923.N13.gz43_252820	F	M00042516B:E03	UC2-ColonMetLiver
3498	451518	1923.A14.gz43_252823	F	M00042352A:G05	UC2-ColonMetLiver
3499	349	1923.M14.gz43_252835	F	M00042460C:B02	UC2-ColonMetLiver
3500	456672	1923.A15.gz43_252839	F	M00042352A:G09	UC2-ColonMetLiver
3501	497101	1923.D15.gz43_252842	F	M00042443D:E01	UC2-ColonMetLiver
3502	488680	1923.H15.gz43_252846	F	M00042452D:C04	UC2-ColonMetLiver
3503	552236	1923.L15.gz43_252850	F	M00042459D:B07	UC2-ColonMetLiver
3504	450823	1923.B17.gz43_252872	F	M00042354B:A07	UC2-ColonMetLiver
3505	446263	1923.D17.gz43_252874	F	M00042444A:C10	UC2-ColonMetLiver
3506	492094	1923.H17.gz43_252878	F	M00042452D:G06	UC2-ColonMetLiver
3507	552581	1923.K17.gz43_252881	F	M00042458C:E06	UC2-ColonMetLiver
3508	494362	1923.G18.gz43_252893	F	M00042450C:B09	UC2-ColonMetLiver
3509	560420	1923.L18.gz43_252898	F	M00042459D:E03	UC2-ColonMetLiver
3510	562399	1923.N18.gz43_252900	F	M00042516C:C04	UC2-ColonMetLiver
3511	487522	1923.O18.gz43_252901	F	M00042518D:A08	UC2-ColonMetLiver
3512	498194	1923.A19.gz43_252903	F	M00042352B:F03	UC2-ColonMetLiver
3513	451126	1923.B19.gz43_252904	F	M00042354C:F04	UC2-ColonMetLiver
3514	450507	1923.E19.gz43_252907	F	M00042446B:G02	UC2-ColonMetLiver
3515	450723	1923.I19.gz43_252911	F	M00042454D:H10	UC2-ColonMetLiver
3516	492981	1923.J19.gz43_252912	F	M00042456C:H07	UC2-ColonMetLiver
3517	451054	1923.F20.gz43_252924	F	M00042448C:C09	UC2-ColonMetLiver
3518	453667	1923.J20.gz43_252928	F	M00042456D:B06	UC2-ColonMetLiver
3519	562588	1923.K20.gz43_252929	F	M00042458D:E06	UC2-ColonMetLiver
3520	527679	1923.L20.gz43_252930	F	M00042459D:G04	UC2-ColonMetLiver
3521	494133	1923.B22.gz43_252952	F	M00042355A:A12	UC2-ColonMetLiver
3522	492304	1923.I22.gz43_252959	F	M00042455A:G12	UC2-ColonMetLiver
3523	100821	1923.J22.gz43_252960	F	M00042456D:D07	UC2-ColonMetLiver
3524	562603	1923.M22.gz43_252963	F	M00042460C:G08	UC2-ColonMetLiver
3525	562275	1923.N22.gz43_252964	F	M00042516D:B11	UC2-ColonMetLiver
3526	451302	1923.A23.gz43_252967	F	M00042352C:G01	UC2-ColonMetLiver
3527	451126	1923.B23.gz43_252968	F	M00042355A:C03	UC2-ColonMetLiver
3528	494130	1923.C23.gz43_252969	F	M00042442A:A12	UC2-ColonMetLiver

Table 2

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NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
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3530	562543	1923.M23.gz43_252979	F	M00042460C:H10	UC2-ColonMetLiver
3531	562398	1923.N23.gz43_252980	F	M00042516D:C01	UC2-ColonMetLiver
3532	497517	1923.P23.gz43_252982	F	M00043296A:E08	UC2-ColonMetLiver
3533	450902	1923.A24.gz43_252983	F	M00042352D:A11	UC2-ColonMetLiver
3534	499178	1923.C24.gz43_252985	F	M00042442A:G04	UC2-ColonMetLiver
3535	450507	1923.E24.gz43_252987	F	M00042446D:F04	UC2-ColonMetLiver
3536	448556	1923.F24.gz43_252988	F	M00042449A:H10	UC2-ColonMetLiver
3537	454575	1923.I24.gz43_252991	F	M00042455B:D05	UC2-ColonMetLiver
3538	275	1923.L24.gz43_252994	F	M00042460B:C06	UC2-ColonMetLiver
3539	452719	1934.C01.gz43_253001	F	M00043360C:E07	UC2-ColonMetLiver
3540	449842	1934.G02.gz43_253021	F	M00043369D:B01	UC2-ColonMetLiver
3541	489506	1934.N02.gz43_253028	F	M00043384D:D05	UC2-ColonMetLiver
3542	453893	1934.P02.gz43_253030	F	M00043388A:D05	UC2-ColonMetLiver
3543	452586	1934.A03.gz43_253031	F	M00043356B:E12	UC2-ColonMetLiver
3544	491177	1934.C03.gz43_253033	F	M00043360C:F11	UC2-ColonMetLiver
3545	496586	1934.M03.gz43_253043	F	M00043382C:D07	UC2-ColonMetLiver
3546	490393	1934.O03.gz43_253045	F	M00043387A:E02	UC2-ColonMetLiver
3547	450756	1934.D04.gz43_253050	F	M00043363B:A11	UC2-ColonMetLiver
3548	456492	1934.I04.gz43_253055	F	M00043373D:G04	UC2-ColonMetLiver
3549	490395	1934.O04.gz43_253061	F	M00043387A:E03	UC2-ColonMetLiver
3550	489275	1934.P04.gz43_253062	F	M00043388B:C02	UC2-ColonMetLiver
3551	489368	1934.C05.gz43_253065	F	M00043360D:D01	UC2-ColonMetLiver
3552	450756	1934.E05.gz43_253067	F	M00043364D:H11	UC2-ColonMetLiver
3553	496084	1934.K05.gz43_253073	F	M00043378D:D12	UC2-ColonMetLiver
3554	450193	1934.L05.gz43_253074	F	M00043380D:E10	UC2-ColonMetLiver
3555	499240	1934.M05.gz43_253075	F	M00043382C:G09	UC2-ColonMetLiver
3556	448230	1934.A06.gz43_253079	F	M00043356C:F03	UC2-ColonMetLiver
3557	455707	1934.H06.gz43_253086	F	M00043372B:B01	UC2-ColonMetLiver
3558	496760	1934.N06.gz43_253092	F	M00043385A:E01	UC2-ColonMetLiver
3559	447429	1934.A07.gz43_253095	F	M00043356C:H07	UC2-ColonMetLiver
3560	450551	1934.G07.gz43_253101	F	M00043369D:G10	UC2-ColonMetLiver
3561	451144	1934.J07.gz43_253104	F	M00043376B:C03	UC2-ColonMetLiver
3562	491402	1934.C08.gz43_253113	F	M00043361B:F02	UC2-ColonMetLiver
3563	449956	1934.E08.gz43_253115	F	M00043365A:C06	UC2-ColonMetLiver
3564	449959	1934.G08.gz43_253117	F	M00043370A:C08	UC2-ColonMetLiver
3565	560213	1934.I08.gz43_253119	F	M00043374A:E05	UC2-ColonMetLiver
3566	450482	1934.J08.gz43_253120	F	M00043376B:F06	UC2-ColonMetLiver
3567	449795	1934.B09.gz43_253128	F	M00043359B:A04	UC2-ColonMetLiver
3568	486856	1934.009.gz43_253141	F	M00043387B:A03	UC2-ColonMetLiver
3569	450080	1934.C10.gz43_253145	F	M00043361D:A02	UC2-ColonMetLiver
3570	510272	1934.I10.gz43_253151	F	M00043374A:G04	UC2-ColonMetLiver

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NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
3571	450583	1934.A11.gz43_253159	F	M00043357A;G04	UC2-ColonMetLiver
3572	450193	1934.B11.gz43_253160	F	M00043359B:D06	UC2-ColonMetLiver
3573	134392	1934.M11.gz43_253171	F	M00043383B:F12	UC2-ColonMetLiver
3574	496234	1934.N11.gz43_253172	F	M00043385C:D06	UC2-ColonMetLiver
3575	487893	1934.A12.gz43_253175	F	M00043357B:B02	UC2-ColonMetLiver
3576	491507	1934.D12.gz43_253178	F	M00043363D:F07	UC2-ColonMetLiver
3577	233814	1934.J12.gz43_253184	F	M00043376D:A12	UC2-ColonMetLiver
3578	450189	1934.A13.gz43_253191	F	M00043357B:D01	UC2-ColonMetLiver
3579	488613	1934.E13.gz43_253195	F	M00043365B:C08	UC2-ColonMetLiver
3580	451011	1934.K13.gz43_253201	F	M00043379C:B09	UC2-ColonMetLiver
3581	450207	1934.G14.gz43_253213	F	M00043370B:D08	UC2-ColonMetLiver
3582	451185	1934.J14.gz43 253216	F	M00043376D:D12	UC2-ColonMetLiver
3583	551380	1934.L14.gz43 253218	F	M00043381B:E10	UC2-ColonMetLiver
3584	495143	1934.M14.gz43 253219	F	M00043383D:C07	UC2-ColonMetLiver
3585	487183	1934.O14.gz43 253221	F	M00043387C:A11	UC2-ColonMetLiver
3586	631526	1934.I15.gz43 253231	F	M00043374D:D07	UC2-ColonMetLiver
3587	449780	1934,E16.gz43 253243	F	M00043365C:A09	UC2-ColonMetLiver
3588	491492	1934.F16.gz43 253244	F	M00043368A:F03	UC2-ColonMetLiver
3589	494099	1934.K16.gz43 253249	F	M00043379D:A05	UC2-ColonMetLiver
3590	450819	1934.N16.gz43 253252	F	M00043386A;A11	UC2-ColonMetLiver
3591	488349	1934.P16.gz43 253254	F	M00043389D:B10	UC2-ColonMetLiver
3592	515423	1934.I17.gz43 253263	F	M00043374D:H09	UC2-ColonMetLiver
3593	490846	1934.C18.gz43 253273	F	M00043362C:E02	UC2-ColonMetLiver
3594	556511	1934.L18.gz43 253282	F	M00043381C;E10	UC2-ColonMetLiver
3595	487437	1934.M18.gz43 253283	F	M00043384B;A04	UC2-ColonMetLiver
3596	491240	1934.E19.gz43 253291	F	M00043365C;F06	UC2-ColonMetLiver
3597	452710	1934.H19.gz43_253294	F	M00043373A:G11	UC2-ColonMetLiver
3598	449959	1934.J20.gz43 253312	F	M00043377C:A11	UC2-ColonMetLiver
3599	449978	1934.P20.gz43 253318	F	M00043390B:C04	UC2-ColonMetLiver
3600	450242	1934,A21.gz43 253319	F	M00043358A:D02	UC2-ColonMetLiver
3601	451294	1934.I21.gz43 253327	F	M00043375A:E02	UC2-ColonMetLiver
3602	450080	1934.G22.gz43 253341	F	M00043371B:C10	UC2-ColonMetLiver
3603	451361	1934,J22.gz43 253344	F	M00043377D;E01	UC2-ColonMetLiver
3604	489207	1934,M22,gz43 253347	F	M00043384C:C02	UC2-ColonMetLiver
3605	488030	1934.O22.gz43 253349	F	M00043387D:B02	UC2-ColonMetLiver
3606	449908	1934.G23.gz43 253357	F	M00043371C:B02	UC2-ColonMetLiver
3607	450211	1934.O23.gz43 253365	F	M00043387D;D06	UC2-ColonMetLiver
3608	402916	1934.I24.gz43 253375	F	M00043375B;A04	UC2-ColonMetLiver
3609	494450	1934,J24.gz43 253376	F	M00043378B;B05	UC2-ColonMetLiver
3610	492627	1934.M24.gz43 253379	F	M00043384C:G01	UC2-ColonMetLiver
3611	550246	1947.J01.gz43 253404	F	M00054555D:C03	UC2-ColonMetLiver
3612	550952	1947,I06.gz43 253483	F	M00054555A:H09	UC2-ColonMetLiver
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Table 2

T	able 2				
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3614	496586	1947.P21.gz43_253730	F	M00054566D:G08	UC2-ColonMetLiver
3615	515707	1947.G22.gz43_253737	F	M00054553D:E09	UC2-ColonMetLiver
3616	552092	1959.F03.gz43 253816	F	M00054631D:C02	UC2-ColonMetLiver
3617	549210	1959.M04.gz43_253839	F	M00054640B:C05	UC2-ColonMetLiver
3618	555923	1959.D08.gz43 253894	F	M00054629C:G06	UC2-ColonMetLiver
3619	461	1959.P08.gz43_253906	F	M00054644B:F02	UC2-ColonMetLiver
3620	551975	1959.D19.gz43_254070	F	M00054630B:A06	UC2-ColonMetLiver
3621	549516	1959.N22.gz43_254128	F	M00054642D:F11	UC2-ColonMetLiver
3622	555949	1959.K24.gz43_254157	F	M00054638D:F02	UC2-ColonMetLiver
3623	193991	1960.L03.gz43_254206	F	M00054661A:B03	UC2-ColonMetLiver
3624	561590	1960.I07.gz43_254267	F	M00054656C:E10	UC2-ColonMetLiver
3625	450867	1960.C12.gz43_254341	F	M00054648C:H10	UC2-ColonMctLiver
3626	264575	1960.N14.gz43_254384	F	M00054664D:A01	UC2-ColonMetLiver
3627	502683	1960.B23.gz43_254516	F	M00054647D:H02	UC2-ColonMetLiver
3628	452662	1960,E24.gz43_254535	F	M00054651D:D02	UC2-ColonMetLiver
3629	551269	1969.F03.gz43_254584	F	M00054674B:B03	UC2-ColonMetLiver
3630	579481	1969.C07.gz43_254645	F	M00054671A:H07	UC2-ColonMetLiver
3631	450624	1969.A10.gz43_254691	F	M00054668A;D01	UC2-ColonMetLiver
3632	415326	1969.N11.gz43_254720	F	M00054685D:D09	UC2-ColonMetLiver
3633	573733	1969.C24.gz43_254917	F	M00054671D:A12	UC2-ColonMetLiver
3634	551659	1983.D01.gz43_254934	F	M00054811D:F01	UC2-ColonMetLiver
3635	549816	1983.E06.gz43_255015	F	M00054812D:H05	UC2-ColonMetLiver
3636	557525	1983.F09.gz43_255064	F	M00054814A:F07	UC2-ColonMetLiver
3637	447405	1983.K11.gz43_255101	F	M00054823B:E07	UC2-ColonMetLiver
3638	494471	1983.C19.gz43_255221	F	M00054811B:H12	UC2-ColonMetLiver
3639	456520	1983.E19.gz43_255223	F	M00054813B:D11	UC2-ColonMetLiver
3640	533520	1984.N02.gz43_255344	F	M00054848B:E02	UC2-ColonMetLiver
3641	560984	1984.P06.gz43_255410	F	M00054851B:E03	UC2-ColonMctLiver
3642	562302	1984.G07.gz43_255417	F	M00054839D:F08	UC2-ColonMetLiver
3643	554196	1984.A12.gz43_255491	F	M00054831A:E11	UC2-ColonMetLiver
3644	557703	1984.G21.gz43_255641	F	M00054840D:B03	UC2-ColonMetLiver
3645	562243	1994.E15.gz43_255927	F	M00054877A:F01	UC2-ColonMetLiver
3646	559531	1994.A19.gz43_255987	F	M00054872A:H10	UC2-ColonMetLiver
3647	447483	1994.F22.gz43_256040	F	M00054878B:G03	UC2-ColonMetLiver
3648	549391	1994.K24.gz43_256077	F	M00054885C:G06	UC2-ColonMetLiver
3649	550204	1995.C02.gz43_256101	F	M00054893D:F07	UC2-ColonMetLiver
3650	554604	1995.K02.gz43_256109	F	M00054904A:F07	UC2-ColonMetLiver
3651	558326	1995.N02.gz43_256112	F	M00054908D:A07	UC2-ColonMetLiver
3652	557218	1995,A03.gz43_256115	F	M00054891D:H05	UC2-ColonMetLiver
3653	562152	1995.C03.gz43_256117	F	M00054893D:G09	UC2-ColonMetLiver
3654	562679	1995.F03.gz43_256120	F	M00054897C:F03	UC2-ColonMetLiver
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3657	549376	1995.K03.gz43_256125	F	M00054904A:H10	UC2-ColonMetLiver
3658	522220	1995,M03,gz43_256127	F	M00054908A:F10	UC2-ColonMetLiver
3659	550710	1995.003.gz43_256129	F	M00054922A:C11	UC2-ColonMetLiver
3660	562672	1995.E04.gz43_256135	F	M00054896B:E09	UC2-ColonMetLiver
3661	559088	1995,I04,gz43_256139	F	M00054901C:A04	UC2-ColonMetLiver
3662	447858	1995.D05.gz43_256150	F	M00054895B;D09	UC2-ColonMetLiver
3663	492256	1995.L05.gz43_256158	F	M00054906D:D06	UC2-ColonMetLiver
3664	550047	1995.B06.gz43_256164	F	M00054893A:F08	UC2-ColonMetLiver
3665	562579	1995.P06.gz43_256178	F	M00054923D:A08	UC2-ColonMetLiver
3666	556804	1995.F07.gz43_256184	F	M00054897D:E08	UC2-ColonMetLiver
3667	553102	1995.I07.gz43_256187	F	M00054901C:G01	UC2-ColonMetLiver
3668	559118	1995.P07.gz43_256194	F	M00054923D:D04	UC2-ColonMetLiver
3669	559554	1995.H08.gz43_256202	F	M00054900B:D12	UC2-ColonMctLiver
3670	510309	1995.K08.gz43_256205	F	M00054904C:B06	UC2-ColonMetLiver
3671	562492	1995.N08.gz43_256208	F	M00054910A:H12	UC2-ColonMetLiver
3672	553975	1995.O08.gz43_256209	F	M00054922B:A08	UC2-ColonMetLiver
3673	561525	1995.N09.gz43_256224	F	M00054910B:F07	UC2-ColonMctLiver
3674	150839	1995.O09.gz43_256225	F	M00054922B:A10	UC2-ColonMetLiver
3675	451049	1995.E10.gz43_256231	F	M00054896C:F01	UC2-ColonMetLiver
3676	560668	1995.H10.gz43_256234	F	M00054900B:G10	UC2-ColonMetLiver
3677	555260	1995.I10.gz43_256235	F	M00054901D:B01	UC2-ColonMetLiver
3678	555785	1995.J10.gz43_256236	F	M00054903C:F03	UC2-ColonMetLiver
3679	82864	1995.D11.gz43_256246	F	M00054895C:B07	UC2-ColonMctLiver
3680	559554	1995.H11.gz43_256250	F	M00054900C:B04	UC2-ColonMetLiver
3681	556404	1995.C12.gz43_256261	F	M00054894C:B11	UC2-ColonMetLiver
3682	516691	1995.H12.gz43_256266	F	M00054900C:C03	UC2-ColonMetLiver
3683	556499	1995.E13.gz43_256279	F	M00054896D:C01	UC2-ColonMctLiver
3684	453715	1995,F13.gz43_256280	F	M00054898A:G09	UC2-ColonMetLiver
3685	556753	1995.H13.gz43_256282	F	M00054900C:D08	UC2-ColonMetLiver
3686	549847	1995.J13.gz43_256284	F	M00054903C:G03	UC2-ColonMetLiver
3687	549822	1995.K13.gz43_256285	F	M00054904C:G09	UC2-ColonMctLiver
3688	489001	1995.L13.gz43_256286	F	M00054907B:E07	UC2-ColonMetLiver
3689	562989	1995.P13.gz43_256290	F	M00054924B:E10	UC2-ColonMetLiver
3690	555998	1995.A14.gz43_256291	F	M00054892C:G05	UC2-ColonMetLiver
3691	562106	1995.B14.gz43_256292	F	M00054893C:A06	UC2-ColonMetLiver
3692	557326	1995.E14.gz43_256295	F	M00054896D;C11	UC2-ColonMetLiver
3693	558935	1995.J14.gz43_256300	F	M00054903C:G04	UC2-ColonMetLiver
3694	399766	1995.L14.gz43_256302	F	M00054907B:F01	UC2-ColonMetLiver
3695	553976	1995.O14.gz43_256305	F	M00054922C:A11	UC2-ColonMetLiver
3696	554002	1995.B15.gz43_256308	F	M00054893C:A07	UC2-ColonMetLiver

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3699	549228	1995.D16.gz43_256326	F	M00054895D:B02	UC2-ColonMetLiver
3700	553534	1995.G16.gz43 256329	F	M00054899C:D11	UC2-ColonMetLiver
3701	557004	1995.H16.gz43_256330	F	M00054900D:F11	UC2-ColonMctLiver
3702	555082	1995,M17.gz43_256351	F	M00054908C:A10	UC2-ColonMetLiver
3703	562714	1995.P17.gz43_256354	F	M00054924D:E01	UC2-ColonMetLiver
3704	552122	1995,D18.gz43_256358	F	M00054895D:C10	UC2-ColonMetLiver
3705	411272	1995.K18.gz43_256365	F	M00054904D:F07	UC2-ColonMctLiver
3706	454812	1995.A19.gz43_256371	F	M00054892D:D06	UC2-ColonMetLiver
3707	558177	1995.J19.gz43_256380	F	M00054903D:G01	UC2-ColonMetLiver
3708	556852	1995.P19.gz43_256386	F	M00054925A:B08	UC2-ColonMetLiver
3709	551463	1995.B20.gz43_256388	F	M00054893C:E04	UC2-ColonMetLiver
3710	558011	1995.G20.gz43_256393	F	M00054899D:C11	UC2-ColonMetLiver
3711	559088	1995.H20.gz43_256394	F	M00054901A:B04	UC2-ColonMetLiver
3712	595066	1995.I20.gz43_256395	F	M00054902B:G02	UC2-ColonMetLiver
3713	558868	1995.K20.gz43_256397	F	M00054904D:G04	UC2-ColonMetLiver
3714	458035	1995.F21.gz43_256408	F	M00054898B:D04	UC2-ColonMetLiver
3715	555837	1995,H21.gz43_256410	F	M00054901A:F01	UC2-ColonMetLiver
3716	452182	1995.I21.gz43_256411	F	M00054902C:A12	UC2-ColonMetLiver
3717	555967	1995.J21.gz43_256412	F	M00054903D:G10	UC2-ColonMetLiver
3718	474163	1995,B22.gz43_256420	F	M00054893C:G06	UC2-ColonMetLiver
3719	556169	1995.H22.gz43_256426	F	M00054901A:H11	UC2-ColonMetLiver
3720	558779	1995.C23.gz43_256437	F	M00054895B:A02	UC2-ColonMetLiver
3721	555830	1995.G23.gz43_256441	F	M00054899D:F07	UC2-ColonMetLiver
3722	974	1995.N23.gz43_256448	F	M00054921C:G01	UC2-ColonMetLiver
3723	556632	1995.B24.gz43_256452	F	M00054893D:D12	UC2-ColonMetLiver
3724	596809	1995.C24.gz43_256453	F	M00054895B:A08	UC2-ColonMctLiver
3725	495365	1995.N24.gz43_256464	F	M00054921C:G06	UC2-ColonMetLiver
3726	447639	1995.024.gz43_256465	F	M00054923A:H02	UC2-ColonMetLiver
3727	418512	1996.K02.gz43_256517	F	M00054938A:B08	UC2-ColonMetLiver
3728	559728	1996.A04.gz43_256539	F	M00054925B:B01	UC2-ColonMetLiver
3729	67060	1996.P05.gz43_256570	F	M00054942C;B04	UC2-ColonMetLiver
3730	549171	1996.E06.gz43 256575	F	M00054931C:F10	UC2-ColonMetLiver
3731	562550	1996.O06.gz43_256585	F	M00054941C:G04	UC2-ColonMetLiver
3732	552566	1996.M08.gz43 256615	F	M00054939D:E03	UC2-ColonMetLiver
3733	473512	1996.B13.gz43_256684	F	M00054927C:C03	UC2-ColonMetLiver
3734	553535	1996.N16.gz43_256744	F	M00054940D:E06	UC2-ColonMetLiver
3735	555739	1996.I18.gz43_256771	F	M00054936B:G02	UC2-ColonMetLiver
3736	873	1996.N18.gz43_256776	F	M00054941A: A08	UC2-ColonMetLiver
3737	549805	1996.I19.gz43 256787	F	M00054936C;B08	UC2-ColonMetLiver
3738	550782	1996.B20.gz43_256796	F	M00054928A;C11	UC2-ColonMetLiver

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3741	447585	1996.I21.gz43 256819	F	M00054936C:G03	UC2-ColonMetLiver
3742	561878	2005.K03.gz43 256919	F	M00054954A:F12	UC2-ColonMetLiver
3743	554833	2005.C07.gz43 256975	F	M00054945B:G09	UC2-ColonMetLiver
3744	126	2005.J08.gz43 256998	F	M00054952D:A11	UC2-ColonMetLiver
3745	496870	2005.I10.gz43 257029	F	M00054952A;B12	UC2-ColonMetLiver
3746	555526	2005.J12.gz43 257062	F	M00054953B:D07	UC2-ColonMetLiver
3747	562053	2005.A14.gz43 257085	F	M00054943D:F12	UC2-ColonMetLiver
3748	450524	2006.B02.gz43 257278	F	M00054961C:H07	UC2-ColonMetLiver
3749	557780	2006.P02.gz43 257292	F	M00054974C:A04	UC2-ColonMetLiver
3750	456795	2006.A03.gz43 257293	F	M00054960C:C03	UC2-ColonMetLiver
3751	556954	2006.E03.gz43 257297	F	M00054964A:F09	UC2-ColonMetLiver
3752	553021	2006.G03.gz43 257299	F	M00054966A:C11	UC2-ColonMetLiver
3753	494306	2006.H03.gz43 257300	F	M00054966D:H12	UC2-ColonMetLiver
3754	208278	2006.I03.gz43 257301	F	M00054967D:F08	UC2-ColonMetLiver
3755	504400	2006.L03.gz43 257304	F	M00054970C:C08	UC2-ColonMetLiver
3756	456203	2006.H05.gz43 257332	F	M00054967A:D11	UC2-ColonMetLiver
3757	599012	2006.C06.gz43 257343	F	M00054962C:D02	UC2-ColonMetLiver
3758	555394	2006.D06.gz43 257344	F	M00054963C:C08	UC2-ColonMetLiver
3759	593173	2006.G06.gz43 257347	F	M00054966A:D09	UC2-ColonMetLiver
3760	592941	2006.I06.gz43 257349	F	M00054968A:C11	UC2-ColonMetLiver
3761	554379	2006.K06.gz43 257351	F	M00054969D:D04	UC2-ColonMetLiver
3762	555103	2006.O06.gz43 257355	F	M00054973B:A10	UC2-ColonMetLiver
3763	556123	2006.J08.gz43 257382	F	M00054969A:E05	UC2-ColonMetLiver
3764	557421	2006.B09.gz43 257390	F	M00054961D:H08	UC2-ColonMetLiver
3765	450932	2006.K09.gz43 257399	F	M00054969D:G11	UC2-ColonMetLiver
3766	553204	2006.E10.gz43 257409	F	M00054964C:B07	UC2-ColonMetLiver
3767	556126	2006.G10.gz43 257411	F	M00054966A:G02	UC2-ColonMetLiver
3768	554204	2006.M11.gz43 257433	F	M00054971C:C11	UC2-ColonMetLiver
3769	556552	2006.P11.gz43_257436	F	M00054974D:C11	UC2-ColonMetLiver
3770	554524	2006.N12.gz43 257450	F	M00054972B:E06	UC2-ColonMetLiver
3771	559047	2006.P12.gz43 257452	F	M00054975A;C08	UC2-ColonMetLiver
3772	554365	2006.P15.gz43_257500	F	M00054975B:B06	UC2-ColonMetLiver
3773	556740	2006,A17,gz43 257517	F	M00054961B:D07	UC2-ColonMetLiver
3774	531145	2006.C17.gz43_257519	F	M00054963A:C12	UC2-ColonMetLiver
3775	557883	2006.F18.gz43 257538	F	M00054965C:C04	UC2-ColonMetLiver
3776	470386	2006.K18.gz43 257543	F	M00054970A;G03	UC2-ColonMetLiver
3777	596152	2006.M18.gz43_257545	F	M00054970A:G05	UC2-ColonMetLiver
3778	556552	2006.J19.gz43_237558	F	M00054969B:F12	UC2-ColonMetLiver
3779	560696	2006.C20.gz43_257567	F	M00054963A:E02	UC2-ColonMetLiver
3780	218113	2006.C20.gz43_257567 2006.K20.gz43_257575	F	M00054963A;E02 M00054970B;B02	UC2-ColonMetLiver
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3782	129715	2006.B22.gz43_257598	F	M00054962A:H07	UC2-ColonMetLiver
3783	553976	2006.C22,gz43_257599	F	M00054963A:G12	UC2-ColonMetLiver
3784	561984	2006.D22.gz43_257600	F	M00054964A:B09	UC2-ColonMetLiver
3785	497938	2006.G22.gz43_257603	F	M00054966D:C05	UC2-ColonMetLiver
3786	556951	2006.J22.gz43_257606	F	M00054969C:G09	UC2-ColonMetLiver
3787	553983	2006.N22.gz43_257610	F	M00054972D:A10	UC2-ColonMetLiver
3788	549579	2006.P22.gz43_257612	F	M00054975C:D08	UC2-ColonMetLiver
3789	555135	2006.B23.gz43_257614	F	M00054962B:A07	UC2-ColonMetLiver
3790	471257	2006.C23.gz43_257615	F	M00054963A:H04	UC2-ColonMetLiver
3791	557080	2006.J23.gz43_257622	F	M00054969C:H05	UC2-ColonMetLiver
3792	554643	2006.M23.gz43_257625	F	M00054971D:F04	UC2-ColonMetLiver
3793	558707	2007.A01.gz43_257645	F	M00054975C:E02	UC2-ColonMetLiver
3794	557420	2007.M02.gz43_257673	F	M00054993D:F04	UC2-ColonMetLiver
3795	482509	2007.A03.gz43_257677	F	M00054975C:G06	UC2-ColonMetLiver
3796	471827	2007.J03.gz43 257686	F	M00054989B:C11	UC2-ColonMetLiver
3797	556841	2007.K03.gz43 257687	F	M00054991B:E11	UC2-ColonMetLiver
3798	597957	2007.L03.gz43 257688	F	M00054992D:C12	UC2-ColonMetLiver
3799	556768	2007.C04.gz43 257695	F	M00054978C:E07	UC2-ColonMetLiver
3800	170450	2007.F04.gz43 257698	F	M00054983C:F05	UC2-ColonMetLiver
3801	555593	2007.I04.gz43 257701	F	M00054987B:D06	UC2-ColonMetLiver
3802	556839	2007.J04.gz43_257702	F	M00054989B:E04	UC2-ColonMetLiver
3803	551967	2007.A05.gz43_257709	F	M00054976A:A03	UC2-ColonMetLiver
3804	550814	2007.I05.gz43_257717	F	M00054987B:G05	UC2-ColonMetLiver
3805	557811	2007.L05.gz43_257720	F	M00054992D:F06	UC2-ColonMetLiver
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3807	554764	2007.N06.gz43 257738	F	M00054995B:G12	UC2-ColonMetLiver
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3809	555639	2007.H07.gz43_257748	F	M00054986B:D07	UC2-ColonMetLiver
3810	556561	2007.E08.gz43_257761	F	M00054981D:C06	UC2-ColonMetLiver
3811	454463	2007.I08.gz43_257765	F	M00054987C:A11	UC2-ColonMetLiver
3812	454463	2007.K08.gz43_257767	F	M00054991C:E01	UC2-ColonMetLiver
3813	562750	2007.C09.gz43_257775	F	M00054978D:H09	UC2-ColonMetLiver
3814	560652	2007.F09.gz43_257778	F	M00054983D:D05	UC2-ColonMetLiver
3815	553615	2007.I09.gz43_257781	F	M00054987C:B12	UC2-ColonMetLiver
3816	555512	2007.F10.gz43_257794	F	M00054984A:A05	UC2-ColonMetLiver
3817	556598	2007.I10.gz43_257797	F	M00054987C:G04	UC2-ColonMetLiver
3818	552188	2007.J10.gz43_257798	F	M00054990A:F10	UC2-ColonMetLiver
3819	558670	2007.A11.gz43_257805	F	M00054976C:A03	UC2-ColonMetLiver
3820	452682	2007.M11.gz43_257817	F	M00054994B:D11	UC2-ColonMetLiver
3821	553915	2007.P11.gz43_257820	F	M00055000A:H10	UC2-ColonMetLiver
3822	550886	2007.C12.gz43 257823	F	M00054979A:H10	UC2-ColonMetLiver

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Table 2	

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3825	554212	2007.O12.gz43_257835	F	M00054997A:C02	UC2-ColonMetLiver
3826	555883	2007.P12.gz43_257836	F	M00055000B:B04	UC2-ColonMetLiver
3827	458257	2007.B13.gz43_257838	F	M00054977C:A09	UC2-ColonMetLiver
3828	555197	2007.D13.gz43_257840	F	M00054980D:H07	UC2-ColonMetLiver
3829	553732	2007.P13.gz43_257852	F	M00055000B:F02	UC2-ColonMetLiver
3830	556416	2007.B14.gz43_257854	F	M00054977C:B08	UC2-ColonMetLiver
3831	501056	2007.B15.gz43_257870	F	M00054977C:D08	UC2-ColonMetLiver
3832	552437	2007.E15.gz43_257873	F	M00054982C:A02	UC2-ColonMetLiver
3833	555658	2007.L15.gz43_257880	F	M00054993B:H06	UC2-ColonMetLiver
3834	476732	2007.O15.gz43_257883	F	M00054997A:G11	UC2-ColonMctLiver
3835	556690	2007.B16.gz43_257886	F	M00054977C:D12	UC2-ColonMetLiver
3836	557615	2007.H16.gz43_257892	F	M00054986D:D11	UC2-ColonMetLiver
3837	543429	2007.G17.gz43_257907	F	M00054985C:B12	UC2-ColonMetLiver
3838	554793	2007.H17.gz43_257908	F	M00054986D:E03	UC2-ColonMetLiver
3839	556396	2007.I17.gz43_257909	F	M00054988C:B08	UC2-ColonMetLiver
3840	556288	2007.J17.gz43_257910	F	M00054990C:A08	UC2-ColonMetLiver
3841	552055	2007.L17.gz43_257912	F	M00054993C:C10	UC2-ColonMetLiver
3842	549516	2007.P17.gz43_257916	F	M00055000C:F08	UC2-ColonMetLiver
3843	561112	2007.B18.gz43_257918	F	M00054977C:F09	UC2-ColonMetLiver
3844	557054	2007.C18.gz43_257919	F	M00054979B:G12	UC2-ColonMetLiver
3845	483410	2007.D18.gz43_257920	F	M00054981B:B09	UC2-ColonMetLiver
3846	598589	2007.G18.gz43_257923	F	M00054985C:D04	UC2-ColonMetLiver
3847	559096	2007.O18.gz43_257931	F	M00054997B:F10	UC2-ColonMetLiver
3848	556759	2007.I19.gz43_257941	F	M00054988C:G02	UC2-ColonMetLiver
3849	555277	2007.K19.gz43_257943	F	M00054992A:C08	UC2-ColonMetLiver
3850	551694	2007.B20.gz43_257950	F	M00054977D:B06	UC2-ColonMetLiver
3851	562655	2007.E20.gz43_257953	F	M00054983 A:F08	UC2-ColonMetLiver
3852	559053	2007.H20.gz43_257956	F	M00054987A:A09	UC2-ColonMetLiver
3853	556286	2007.J20.gz43_257958	F	M00054990D:A06	UC2-ColonMetLiver
3854	551798	2007.K20.gz43_257959	F	M00054992A:D11	UC2-ColonMetLiver
3855	549214	2007.N20.gz43_257962	F	M00054996A:D10	UC2-ColonMetLiver
3856	467293	2007.D21.gz43_257968	F	M00054981B:H12	UC2-ColonMetLiver
3857	503452	2007.J21.gz43_257974	F	M00054990D:F04	UC2-ColonMetLiver
3858	556019	2007.L22.gz43_257992	F	M00054993C:G12	UC2-ColonMetLiver
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3860	562323	2007.L23.gz43_258008	F	M00054993C:H05	UC2-ColonMetLiver
3861	2284	2007.A24.gz43_258013	F	M00054976D:F06	UC2-ColonMetLiver
3862	481243	2007.D24.gz43_258016	F	M00054981C:A11	UC2-ColonMetLiver
3863	555958	2007.N24.gz43_258026	F	M00054996B:C11	UC2-ColonMetLiver
3864	560984	2007.O24.gz43_258027	F	M00054997C:H03	UC2-ColonMetLiver

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3867	553064	2008.A02.gz43_258047	F	M00055001A:B10	UC2-ColonMetLiver
3868	552006	2008.D02.gz43 258050	F	M00055005A;C07	UC2-ColonMetLiver
3869	559676	2008.E02.gz43 258051	F	M00055006A:D01	UC2-ColonMetLiver
3870	557308	2008.J02.gz43 258056	F	M00055011B:F09	UC2-ColonMetLiver
3871	549723	2008.N02.gz43_258060	F	M00055016D:C11	UC2-ColonMetLiver
3872	549912	2008.B03.gz43_258064	F	M00055002B:E07	UC2-ColonMetLiver
3873	551444	2008.G03.gz43_258069	F	M00055008C:F09	UC2-ColonMetLiver
3874	557240	2008.P03.gz43_258078	F	M00055019B:H10	UC2-ColonMetLiver
3875	559011	2008.C04.gz43 258081	F	M00055003C;E08	UC2-ColonMetLiver
3876	555701	2008.D04.gz43_258082	F	M00055005A:E02	UC2-ColonMetLiver
3877	556115	2008.P04.gz43 258094	F	M00055019C:C01	UC2-ColonMetLiver
3878	556042	2008.B05.gz43 258096	F	M00055002B:G03	UC2-ColonMetLiver
3879	558212	2008.E05.gz43 258099	F	M00055006B:A09	UC2-ColonMetLiver
3880	554932	2008.G05.gz43_258101	F	M00055008C:H10	UC2-ColonMetLiver
3881	548858	2008.J05.gz43 258104	F	M00055011C:E04	UC2-ColonMetLiver
3882	605117	2008.L05.gz43_258106	F	M00055014B:E08	UC2-ColonMetLiver
3883	560525	2008,M05.gz43 258107	F	M00055015D:C09	UC2-ColonMetLiver
3884	597780	2008.I06.gz43 258119	F	M00055010C:C10	UC2-ColonMetLiver
3885	411113	2008.N06.gz43 258124	F	M00055017A:A11	UC2-ColonMetLiver
3886	552674	2008.E07.gz43 258131	F	M00055006B:C08	UC2-ColonMetLiver
3887	495942	2008.O07.gz43 258141	F	M00055018C:C06	UC2-ColonMetLiver
3888	34381	2008.F08.gz43 258148	F	M00055007C:D01	UC2-ColonMetLiver
3889	549195	2008.H08.gz43_258150	F	M00055009C:F06	UC2-ColonMetLiver
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3891	557121	2008.P08.gz43 258158	F	M00055019C:G06	UC2-ColonMetLiver
3892	554693	2008.A09.gz43 258159	F	M00055001A:G05	UC2-ColonMetLiver
3893	134237	2008.C09.gz43 258161	F	M00055003D:F09	UC2-ColonMetLiver
3894	594994	2008.H09.gz43 258166	F	M00055009C:F07	UC2-ColonMetLiver
3895	555371	2008.I09.gz43 258167	F	M00055010D:A02	UC2-ColonMetLiver
3896	347057	2008.E10.gz43_258179	F	M00055006B;E05	UC2-ColonMetLiver
3897	558378	2008.H10.gz43 258182	F	M00055009D:D12	UC2-ColonMetLiver
3898	497086	2008.K10.gz43 258185	F	M00055013B:B07	UC2-ColonMetLiver
3899	554379	2008.B11.gz43_258192	F	M00055002D:A07	UC2-ColonMetLiver
3900	451118	2008.L11.gz43 258202	F	M00055014D:A11	UC2-ColonMetLiver
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3904	562550	2008.B12.gz43_258208	F	M00055002D:E04	UC2-ColonMetLiver
3905	556530	2008.L12.gz43_258218	F	M00055015A:C08	UC2-ColonMetLiver
3906	524897	2008.O12.gz43_258221	F	M00055018D:E05	UC2-ColonMetLiver

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3989 549114 2017.B22.gz43_258754 F M00055021D:F06 UC2-ColonMetLiver	3987	529356		F	M00055024B:E06	
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3990 556933 2017.D22.gz43 258756 F M00055024B:F02 UC2-ColonMetLiver	3989	549114	2017.B22.gz43_258754	F	M00055021D:F06	UC2-ColonMetLiver
	3990	556933	2017.D22.gz43 258756	F	M00055024B:F02	UC2-ColonMetLiver

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Table 2

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3993	460078	2017.L23.gz43_258780	F	M00055034A:G01	UC2-ColonMetLiver
3994	561836	2017.N23.gz43_258782	F	M00055036B:H02	UC2-ColonMetLiver
3995	558185	2043.D03.gz43_259228	F	M00055222D:B11	UC2-ColonMetLiver
3996	550632	2043.F03.gz43_259230	F	M00055225C:E08	UC2-ColonMetLiver
3997	551677	2043.J03.gz43_259234	F	M00055230A:H01	UC2-ColonMetLiver
3998	451720	2043.N03.gz43_259238	F	M00055236C:E05	UC2-ColonMctLiver
3999	559447	2043.L05.gz43_259268	F	M00055233C:F09	UC2-ColonMctLiver
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4001	554627	2043.A06.gz43_259273	F	M00055219B:B04	UC2-ColonMetLiver
4002	558422	2043.N06.gz43_259286	F	M00055236D:E04	UC2-ColonMetLiver
4003	558679	2043.G08.gz43_259311	F	M00055226C:H05	UC2-ColonMetLiver
4004	491240	2043.F09.gz43_259326	F	M00055225D:G11	UC2-ColonMetLiver
4005	550815	2043.H09.gz43_259328	F	M00055227C:D02	UC2-ColonMetLiver
4006	481362	2043.P10.gz43_259352	F	M00055239C:F08	UC2-ColonMetLiver
4007	467710	2043.C11.gz43_259355	F	M00055221D:H08	UC2-ColonMetLiver
4008	557965	2043.B12.gz43_259370	F	M00055220D:G07	UC2-ColonMetLiver
4009	380127	2043.C12.gz43_259371	F	M00055222A:A06	UC2-ColonMetLiver
4010	374281	2043.M12.gz43_259381	F	M00055235B:C01	UC2-ColonMetLiver
4011	464067	2043.C13.gz43_259387	F	M00055222A:C08	UC2-ColonMetLiver
4012	552521	2043.D13.gz43_259388	F	M00055223B:C04	UC2-ColonMetLiver
4013	552019	2043.P13.gz43_259400	F	M00055239D:C01	UC2-ColonMetLiver
4014	556790	2043.A14.gz43_259401	F	M00055219C:C10	UC2-ColonMetLiver
4015	557411	2043.K14.gz43_259411	F	M00055232C:D04	UC2-ColonMetLiver
4016	378459	2043.L15.gz43_259428	F	M00055233D:G11	UC2-ColonMetLiver
4017	550571	2043.H16.gz43_259440	F	M00055227D:E07	UC2-ColonMetLiver
4018	557676	2043.P16.gz43_259448	F	M00055239D:G04	UC2-ColonMetLiver
4019	558652	2043.A17.gz43_259449	F	M00055219C:H06	UC2-ColonMctLiver
4020	559053	2043.I17.gz43_259457	F	M00055229C:D01	UC2-ColonMetLiver
4021	552086	2043.C18.gz43_259467	F	M00055222B:A01	UC2-ColonMetLiver
4022	552430	2043.L18.gz43_259476	F	M00055234A;G12	UC2-ColonMetLiver
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4027	557656	2043.O23.gz43_259559	F	M00055238D:D10	UC2-ColonMetLiver
4028	559764	2043.D24.gz43_259564	F	M00055223D:H03	UC2-ColonMetLiver
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4030	552249	2043.P24.gz43_259576	F	M00055240B;E12	UC2-ColonMetLiver
4031	556829	2044.G02.gz43_259603	F	M00055248D:B12	UC2-ColonMetLiver
4032	558093	2044.A03.gz43_259613	F	M00055240C;A03	UC2-ColonMetLiver

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4036	607430	2044.F04.gz43_259634	F	M00055247B:A11	UC2-ColonMetLiver
4037	558512	2044.L04.gz43_259640	F	M00055255B:F05	UC2-ColonMetLiver
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4048	460049	2044.P08.gz43 259708	F	M00055260B:H06	UC2-ColonMetLiver
4049	560317	2044.D09.gz43 259712	F	M00055244D:C07	UC2-ColonMetLiver
4050	560317	2044.E09.gz43 259713	F	M00055246A:C01	UC2-ColonMetLiver
4051	552357	2044.H09.gz43 259716	F	M00055250B:G09	UC2-ColonMetLiver
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4054	423884	2044.N10.gz43_259738	F	M00055258A:C07	UC2-ColonMetLiver
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4071	238121	2044.C20.gz43 259887	F	M00055244A;C06	UC2-ColonMetLiver
4072	629002	2044.D20,gz43 259888	F	M00055245B:H01	UC2-ColonMetLiver
4073	562256	2044.L21.gz43_259912	F	M00055256B:C07	UC2-ColonMetLiver
4074	541784	2044.B22.gz43 259918	F	M00055242D:E02	UC2-ColonMctLiver

Table 2

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4091 556019 2053.D08.gz43 2600089 F M00055264B:E06 UC2-ColonMetLiver 4092 480960 2053.O09.gz43 260110 F M00055279B:D02 UC2-ColonMetLiver 4093 478087 2953.C10.gz43 260120 F M00055269B:D02 UC2-ColonMetLiver 4094 450507 2053.D10.gz43 260121 F M00055265C:D80 UC2-ColonMetLiver 4096 58052 2053.B11.gz43 260135 F M0005526C:D808 UC2-ColonMetLiver 4096 58052 2053.B11.gz43 260150 F M00055261B:G12 UC2-ColonMetLiver 4097 550863 2053.A12.gz43 260161 F M00055274D:A11 UC2-ColonMetLiver 4098 560957 2053.L12.gz43 260161 F M00055274D:A11 UC2-ColonMetLiver 4100 450242 2053.C13.gz43 260168 F M00055265C:F01 UC2-ColonMetLiver 4101 46557 2053.B14.gz43 260187 F M00055265C:F01 UC2-ColonMetLiver </td <td>4089</td> <td>555564</td> <td>2053.N05.gz43_260051</td> <td>F</td> <td>M00055277D:A02</td> <td>UC2-ColonMetLiver</td>	4089	555564	2053.N05.gz43_260051	F	M00055277D:A02	UC2-ColonMetLiver
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	4116	558513	2053.P21.gz43_260309	F	M00055281A:F08	UC2-ColonMetLiver

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4138	450883	1669.E08.gz43 260803	F	M00055657A:B04	UC2-NormColon
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4142	644971	1669.B10.gz43 260832	F	M00055653B:G01	UC2-NormColon
4143		1669.C10.gz43_260833	F	M00055654C:D03	UC2-NormColon
4144	452687	1669.G10.gz43_260837	F	M00055660A:A06	UC2-NormColon
4145	643909	1669.J10.gz43 260840	F	M00055663D:B05	UC2-NormColon
4146	646894	1669.M10.gz43_260843	F	M00055667C:F07	UC2-NormColon
4147	556750	1669.N10.gz43 260844	F	M00055668D:E11	UC2-NormColon
4148	640259	1669.B11.gz43 260848	F	M00055653B:G11	UC2-NormColon
4149	643804	1669.D11.gz43_260850	F	M00055656A:E09	UC2-NormColon
4150	97909	1669.F11.gz43_260852	F	M00055659A:A08	UC2-NormColon
4151	503275	1669.G11.gz43_260853	F	M00055660A:C05	UC2-NormColon
4152	640678	1669.L11.gz43_260858	F	M00055666B:E06	UC2-NormColon
4153	406734	1669.A12.gz43 260863	F	M00055652B:F12	UC2-NormColon
4154	643563	1669.G12.gz43 260869	F	M00055660A:C08	UC2-NormColon
4155	658447	1669.H12.gz43 260870	F	M00055661C:E11	UC2-NormColon
4156	446188	1669.H12.gz43_260870	F	M00055662C:H06	UC2-NormColon UC2-NormColon
4157	603857	1669.L12.gz43_260874	F	M00055662C:H06	UC2-NormColon UC2-NormColon
4158	561894	1669.M12.gz43 260874	F		UC2-NormColon UC2-NormColon
4138	201024	1005.W112.gz45_200875	r	M00055667D:B01	UC2-NormColon

Table 2

	able 2				
SEQ					
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	CLUSTER		NT	CLONE ID	LIBRARY
4159	639596	1669.I15.gz43_260919	F	M00055662D:A09	UC2-NormColon
4160		1669.G17.gz43_260949	F	M00055660B:H02	UC2-NormColon
4161	639665	1669.I17.gz43_260951	F	M00055662D:B07	UC2-NormColon
4162		1669.O17.gz43_260957	F	M00055670D:F02	UC2-NormColon
4163	642111	1669.C18.gz43_260961	F	M00055655A;A09	UC2-NormColon
4164	640179	1669.K18.gz43_260969	F	M00055665D:F11	UC2-NormColon
4165	648352	1669.N18.gz43_260972	F	M00055669B:G02	UC2-NormColon
4166	640018	1669.I20.gz43_260999	F	M00055662D:E05	UC2-NormColon
4167	642332	1669.L20.gz43_261002	F	M00055667A:B12	UC2-NormColon
4168	639706	1669.F21.gz43_261012	F	M00055659C:B10	UC2-NormColon
4169	642411	1669.C22.gz43_261025	F	M00055655B:B08	UC2-NormColon
4170	641091	1669.I22.gz43_261031	F	M00055663A:B02	UC2-NormColon
4171	463312	1669.B23.gz43_261040	F	M00055653D:H02	UC2-NormColon
4172	456561	1669.L23.gz43_261050	F	M00055667A:H10	UC2-NormColon
4173	649631	1669.A24.gz43_261055	F	M00055653A:G08	UC2-NormColon
4174	639674	1669.F24.gz43_261060	F	M00055659C:D06	UC2-NormColon
4175	640261	1669.I24.gz43_261063	F	M00055663A:G04	UC2-NormColon
4176	640889	1670.N03.gz43_261116	F	M00055691B:E07	UC2-NormColon
4177	426366	1670.D05.gz43 261138	F	M00055676D:H07	UC2-NormColon
4178	643594	1670.L05.gz43 261146	F	M00055687C:F01	UC2-NormColon
4179	446675	1670.M05.gz43 261147	F	M00055689B:F04	UC2-NormColon
4180	640645	1670.H06.gz43 261158	F	M00055683B;D10	UC2-NormColon
4181	645909	1670.J06.gz43 261160	F	M00055685B:E12	UC2-NormColon
4182	551287	1670.K06.gz43 261161	F	M00055686B;D03	UC2-NormColon
4183	646638	1670.N06.gz43 261164	F	M00055691C:E02	UC2-NormColon
4184	639805	1670.B07.gz43 261168	F	M00055673D:C01	UC2-NormColon
4185	640092	1670.A08.gz43 261183	F	M00055672D:F06	UC2-NormColon
4186		1670.M08.gz43 261195	F	M00055689C:B03	UC2-NormColon
4187		1670.G09.gz43 261205	F	M00055682C:D06	UC2-NormColon
4188		1670.L09.gz43_261210	F	M00055688A:A02	UC2-NormColon
4189		1670.H10.gz43 261222	F	M00055683C;A03	UC2-NormColon
4190		1670.I10.gz43 261223	F	M00055684B;D06	UC2-NormColon
4191	634122	1670.M10.gz43_261227	F	M00055689C;C03	UC2-NormColon
4192		1670.K12.gz43 261257	F	M00055686D:E04	UC2-NormColon
4193		1670.L12.gz43_261258	F	M00055688A:E04	UC2-NormColon
4194		1670.P12.gz43_261262	F	M00055695D:E01	UC2-NormColon
4195		1670.K13.gz43 261273	F	M00055686D:E10	UC2-NormColon
4196		1670.N13.gz43 261276	F	M00055691D:E04	UC2-NormColon
4197		1670.C14.gz43_261281	F	M00055675D:E04	UC2-NormColon
4198		1670.H14.gz43_261286	F	M00055683C:D11	UC2-NormColon
4199		1670.A16.gz43 261311	F	M00055673B:A04	UC2-NormColon
4200		1670,E17,gz43 261331	F	M00055680B:H04	UC2-NormColon
					_ CD 1101

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Table 2

T	able 2				•
SEQ					
ID			ORIE		
NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
4201	553921	1670.H17.gz43_261334	F	M00055683C:H11	UC2-NormColon
4202	639469	1670.A18.gz43_261343	F	M00055673B:B08	UC2-NormColon
4203	359500	1670.F18.gz43_261348	F	M00055681D:F11	UC2-NormColon
4204	649558	1670.J18.gz43_261352	F	M00055685D:B06	UC2-NormColon
4205	561920	1670.N18.gz43_261356	F	M00055692A:E05	UC2-NormColon
4206	446242	1670.C20.gz43_261377	F	M00055676B:D05	UC2-NormColon
4207	645968	1670.B21.gz43_261392	F	M00055674C:E05	UC2-NormColon
4208	637966	1670.F22.gz43_261412	F	M00055682A:B07	UC2-NormColon
4209	640826	1670.M22.gz43_261419	F	M00055691A:D08	UC2-NormColon
4210	642321	1670.D23.gz43_261426	F	M00055679B:E03	UC2-NormColon
4211	601137	1670.I23.gz43 261431	F	M00055685A:E10	UC2-NormColon
4212	416808	1670.E24.gz43_261443	F	M00055681A:F02	UC2-NormColon
4213	647856	1670.I24.gz43 261447	F	M00055685A:F02	UC2-NormColon
4214	554564	1670.K24.gz43 261449	F	M00055687C:B04	UC2-NormColon
4215	561422	1671.I07.gz43 261559	F	M00055711B:B08	UC2-NormColon
4216	642288	1671.P07.gz43 261566	F	M00055721B:D08	UC2-NormColon
4217	642062	1671.E08.gz43 261571	F	M00055704A:D09	UC2-NormColon
4218	463368	1671.K08.gz43 261577	F	M00055715A:D10	UC2-NormColon
4219	646028	1671.D09.gz43 261586	F	M00055702C:D01	UC2-NormColon
4220	478192	1671.M09.gz43 261595	F	M00055718A:F01	UC2-NormColon
4221	640947	1671.D10.gz43 261602	F	M00055702C:F11	UC2-NormColon
4222	463548	1671.G10.gz43 261605	F	M00055706B:H12	UC2-NormColon
4223	640282	1671.M10.gz43 261611	F	M00055718A:H05	UC2-NormColon
4224	41141	1671.B11.gz43 261616	F	M00055699C:D02	UC2-NormColon
4225	463368	1671.I11.gz43 261623	F	M00055711C:A07	UC2-NormColon
4226	558086	1671.J11.gz43 261624	F	M00055713C:F12	UC2-NormColon
4227	560546	1671.C12.gz43 261633	F	M00055701A:B04	UC2-NormColon
4228	551415	1671.E12.gz43 261635	F	M00055704B:C04	UC2-NormColon
4229	557710	1671.N12.gz43 261644	F	M00055719A;D11	UC2-NormColon
4230	649429	1671.G14.gz43 261669	F	M00055707A:B07	UC2-NormColon
4231	451401	1671.L14.gz43 261674	F	M00055717B:A03	UC2-NormColon
4232	642039	1671.C17.gz43 261713	F	M00055701B:C03	UC2-NormColon
4233	471268	1671.F17.gz43 261716	F	M00055705C;F12	UC2-NormColon
4234	417259	1671.H17.gz43 261718	F	M00055709D:G10	UC2-NormColon
4235	421826	1671.L17.gz43 261722	F	M00055717B:E04	UC2-NormColon
4236	465589	1671.M17.gz43 261723	F	M00055718B:H11	UC2-NormColon
4237	645505	1671.N17.gz43 261724	F	M00055719A:G04	UC2-NormColon
4238	510195	1671.N18.gz43 261740	F	M00055719A:G12	UC2-NormColon
4239	492094	1671.P18.gz43 261742	F	M00055722A:B05	UC2-NormColon
4240	453006	1671.G20.gz43 261765	F	M00055707A:F11	UC2-NormColon
4241	600115	1671.P21.gz43 261790	F	M00055722A:C04	UC2-NormColon
4242	641700	1671.H22.gz43 261798	F	M00055710B:E04	UC2-NormColon
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Table 2

	able 2				
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NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
4243	466265	1671.O22.gz43_261805	F	M00055721A:A07	UC2-NormColon
4244	553985	1671.B23.gz43_261808	F	M00055700B:E10	UC2-NormColon
4245	536415	1671.L23.gz43_261818	F	M00055717C:B07	UC2-NormColon
4246	640282	1671.N23.gz43_261820	F	M00055719C:A09	UC2-NormColon
4247	498194	1672.A01.gz43_261841	F	M00055722B:A01	UC2-NormColon
4248	641305	1672.E02.gz43_261861	F	M00055725A:F12	UC2-NormColon
4249	641070	1672.H02.gz43_261864	F	M00055727D:G01	UC2-NormColon
4250	556326	1672.B03.gz43_261874	F	M00055723A:B08	UC2-NormColon
4251	522497	1672.C03.gz43_261875	F	M00055723D:C04	UC2-NormColon
4252	557833	1672.E03.gz43_261877	F	M00055725A:G07	UC2-NormColon
4253	647688	1672.B04.gz43_261890	F	M00055723A:C04	UC2-NormColon
4254	416377	1672.M04.gz43_261901	F	M00055733D:H01	UC2-NormColon
4255	643897	1672.004.gz43_261903	F	M00055736B:C08	UC2-NormColon
4256	645171	1672.A06.gz43_261921	F	M00055722B:G12	UC2-NormColon
4257	649106	1672.C06.gz43_261923	F	M00055723D:E05	UC2-NormColon
4258	454383	1672.K06.gz43_261931	F	M00055731C:F09	UC2-NormColon
4259	649360	1672.B07.gz43_261938	F	M00055723A:F04	UC2-NormColon
4260	639458	1672.K08.gz43_261963	F	M00055731C:H07	UC2-NormColon
4261	642078	1672.B11.gz43_262002	F	M00055723B:A09	UC2-NormColon
4262	642631	1672.L11.gz43_262012	F	M00055733A:G04	UC2-NormColon
4263	650303	1672.A12.gz43_262017	F	M00055722C:E11	UC2-NormColon
4264	284586	1672.C12.gz43_262019	F	M00055724A:C12	UC2-NormColon
4265	642535	1672.G12.gz43_262023	F	M00055727B:F10	UC2-NormColon
4266	642491	1672.A13.gz43_262033	F	M00055722C:F11	UC2-NormColon
4267	100821	1672.P13.gz43_262048	F	M00055738B:H11	UC2-NormColon
4268	641056	1672.A14.gz43_262049	F	M00055722D:A07	UC2-NormColon
4269	642197	1672.B14.gz43_262050	F	M00055723B:C03	UC2-NormColon
4270	801	1672.E14.gz43_262053	F	M00055725D:B02	UC2-NormColon
4271	449956	1672.N14.gz43_262062	F	M00055735C:G08	UC2-NormColon
4272	645018	1672.C15.gz43_262067	F	M00055724A:E03	UC2-NormColon
4273	640879	1672.K15.gz43_262075	F	M00055731D:F09	UC2-NormColon
4274	642263	1672.015.gz43_262079	F	M00055736D:D11	UC2-NormColon
4275	647906	1672.P15.gz43_262080	F	M00055738C:C02	UC2-NormColon
4276	559776	1672.G16.gz43_262087	F	M00055727C:B02	UC2-NormColon
4277	647060	1672.H16.gz43_262088	F	M00055728B:G11	UC2-NormColon
4278	642315	1672.I16,gz43_262089	F	M00055729C:D11	UC2-NormColon
4279	504501	1672.L16.gz43_262092	F	M00055733B:F11	UC2-NormColon
4280	460967	1672.A17.gz43_262097	F	M00055722D:B10	UC2-NormColon
4281	504568	1672.C17.gz43_262099	F	M00055724A:G08	UC2-NormColon
4282	559828	1672.D17.gz43_262100	F	M00055724D:A02	UC2-NormColon
4283	455113	1672.H17.gz43_262104	F	M00055728C:B08	UC2-NormColon
4284	477064	1672.I17.gz43_262105	F	M00055729C:E02	UC2-NormColon

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T	able 2				
SEQ					
ID			ORIE		
NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
4285	1558	1672,J17.gz43_262106	F	M00055731A:G05	UC2-NormColon
4286	419443	1672.L17.gz43_262108	F	M00055733B:H05	UC2-NormColon
4287	546705	1672.I18.gz43_262121	F	M00055729D:A06	UC2-NormColon
4288	235397	1672.J18.gz43_262122	F	M00055731A:H07	UC2-NormColon
4289	651085	1672.K18.gz43_262123	F	M00055732A;B04	UC2-NormColon
4290	502826	1672.N18.gz43_262126	F	M00055735D:C01	UC2-NormColon
4291	544797	1672.A19.gz43_262129	F	M00055722D:E05	UC2-NormColon
4292	489426	1672.B19.gz43_262130	F	M00055723C:B02	UC2-NormColon
4293	452212	1672.E19.gz43 262133	F	M00055725D:F05	UC2-NormColon
4294	646609	1672,M19.gz43 262141	F	M00055734D:D01	UC2-NormColon
4295	556336	1672.N19.gz43 262142	F	M00055735D:E10	UC2-NormColon
4296	467822	1672.C20.gz43 262147	F	M00055724B:D04	UC2-NormColon
4297	650397	1672.L20.gz43 262156	F	M00055733C:H12	UC2-NormColon
4298	37862	1672.P20.gz43 262160	F	M00055738D:G08	UC2-NormColon
4299	644609	1672.I21.gz43 262169	F	M00055729D:F05	UC2-NormColon
4300	561968	1672.L21.gz43 262172	F	M00055733D:B11	UC2-NormColon
4301	550049	1672.P21.gz43_262176	F	M00055739A:B02	UC2-NormColon
4302	554708	1672.H22.gz43 262184	F	M00055728D:F02	UC2-NormColon
4303	608540	1672.P22.gz43 262192	F	M00055739A;B04	UC2-NormColon
4304	470769	1672.A23.gz43_262193	F	M00055722D:G12	UC2-NormColon
4305	642722	1672.C23.gz43_262195	F	M00055724B:G03	UC2-NormColon
4306	645151	1672.L23.gz43_262204	F	M00055733D:D04	UC2-NormColon
4307	648265	1672.N23.gz43_262206	F	M00055736A:B11	UC2-NormColon
4308	489249	1672.F24.gz43_262214	F	M00055726D:H12	UC2-NormColon
4309	644173	1682.H02.gz43_262248	F	M00055770A:G08	UC2-NormColon
4310	642417	1682.L02.gz43_262252	F	M00055774D:A05	UC2-NormColon
4311	553877	1682.M02.gz43_262253	F	M00055775C:B10	UC2-NormColon
4312	530238	1682.H03.gz43_262264	F	M00055770A:H11	UC2-NormColon
4313	553380	1682.L03.gz43_262268	F	M00055774D:B07	UC2-NormColon
4314	637387	1682.E04.gz43_262277	F	M00055766B:E11	UC2-NormColon
4315	634660	1682.F04.gz43 262278	F	M00055767A:D10	UC2-NormColon
4316	642791	1682.J04.gz43_262282	F	M00055772A:H08	UC2-NormColon
4317	446163	1682.K04.gz43_262283	F	M00055773C:C09	UC2-NormColon
4318	641056	1682.M04.gz43_262285	F	M00055775C:E10	UC2-NormColon
4319	550562	1682.B05.gz43_262290	F	M00055763B:D07	UC2-NormColon
4320	648320	1682.A06.gz43_262305	F	M00055761D:C03	UC2-NormColon
4321	446757	1682.H06.gz43_262312	F	M00055770B:D06	UC2-NormColon
4322	454540	1682.K06.gz43_262315	F	M00055773C:H12	UC2-NormColon
4323	458979	1682.L06.gz43_262316	F	M00055774D:E02	UC2-NormColon
4324	641174	1682,P06.gz43_262320	F	M00055780C:E02	UC2-NormColon
4325	642852	1682.E08.gz43_262341	F	M00055766B:H10	UC2-NormColon
4326	488680	1682,A09,gz43 262353	F	M00055761D:H08	UC2-NormColon

Table 2

	able 2				
SEQ					
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NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
4327	27083	1682.K09.gz43_262363	F	M00055773D:G11	UC2-NormColon
4328	524470	1682.C10.gz43_262371	F	M00055765A:A04	UC2-NormColon
4329	463951	1682.J10.gz43_262378	F	M00055772C:E11	UC2-NormColon
4330	584179	1682.C11.gz43_262387	F	M00055765A:B02	UC2-NormColon
4331	642460	1682.D11.gz43_262388	F	M00055765D:E06	UC2-NormColon
4332	448944	1682.E11.gz43_262389	F	M00055766C:C11	UC2-NormColon
4333	642082	1682.F11.gz43_262390	F	M00055767B:D02	UC2-NormColon
4334	642095	1682.N11.gz43_262398	F	M00055778D:A03	UC2-NormColon
4335	562876	1682.H12.gz43_262408	F	M00055770C:H11	UC2-NormColon
4336	453079	1682.I12.gz43_262409	F	M00055771C:A11	UC2-NormColon
4337	101499	1682.L12.gz43_262412	F	M00055775A:G09	UC2-NormColon
4338	639829	1682.N12.gz43_262414	F	M00055778D:C03	UC2-NormColon
4339	642332	1682.P12.gz43_262416	F	M00055780D:D09	UC2-NormColon
4340	541499	1682.G13.gz43_262423	F	M00055768B:H12	UC2-NormColon
4341	509410	1682.I13.gz43_262425	F	M00055771C:D09	UC2-NormColon
4342	89082	1682.O13.gz43_262431	F	M00055780A:C04	UC2-NormColon
4343	453091	1682.P13.gz43_262432	F	M00055780D:F08	UC2-NormColon
4344	639981	1682.A14.gz43_262433	F	M00055762B:B11	UC2-NormColon
4345	642564	1682.B14.gz43_262434	F	M00055763C:G08	UC2-NormColon
4346	561558	1682.O14.gz43_262447	F	M00055780A:C06	UC2-NormColon
4347	562870	1682.B15.gz43_262450	F	M00055763D:B11	UC2-NormColon
4348	462742	1682.D15.gz43_262452	F	M00055766A:A01	UC2-NormColon
4349	451037	1682.E15.gz43_262453	F	M00055766C:G12	UC2-NormColon
4350	641563	1682.J15.gz43_262458	F	M00055772D:C10	UC2-NormColon
4351	643940	1682.A16.gz43_262465	F	M00055762C:B04	UC2-NormColon
4352	642204	1682.C16.gz43_262467	F	'M00055765A:C11	UC2-NormColon
4353	517912	1682.J16.gz43_262474	F	M00055772D:D03	UC2-NormColon
4354	641576	1682.D17.gz43_262484	F	M00055766A:D08	UC2-NormColon
4355	642082	1682.F17.gz43_262486	F	M00055767D:A12	UC2-NormColon
4356	450211	1682.O17.gz43_262495	F	M00055780A:E11	UC2-NormColon
4357	621636	1682.B18.gz43_262498	_ F	M00055764A:F02	UC2-NormColon
4358	639232	1682.D18.gz43_262500	F	M00055766A:E08	UC2-NormColon
4359	450553	1682.J18.gz43_262506	F	M00055772D:F10	UC2-NormColon
4360	647940	1682.K18.gz43_262507	F	M00055774B:F07	UC2-NormColon
4361	461835	1682.M18.gz43_262509	F	M00055778A;F09	UC2-NormColon
4362	641496	1682.O18.gz43_262511	F	M00055780A:F07	UC2-NormColon
4363	464510	1682.H19.gz43_262520	F	M00055770D:E10	UC2-NormColon
4364	642564	1682.M19.gz43_262525	F	M00055778A:F12	UC2-NormColon
4365	629002	1682.C20.gz43_262531	. F	M00055765A:G02	UC2-NormColon
4366	1093	1682.D20.gz43_262532	F	M00055766A:H03	UC2-NormColon
4367	50351	1682.E20.gz43_262533	F	M00055766D:D05	UC2-NormColon
4368	642417	1682.F20.gz43_262534	F	M00055767D:E07	UC2-NormColon

Table 2

	able 2				
SEQ					
ID			ORIE		
NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
4369	466971	1682.J20.gz43_262538	F	M00055772D:H04	UC2-NormColon
4370	641185	1682.K20.gz43_262539	F	M00055774B:H01	UC2-NormColon
4371	644240	1682.L20.gz43_262540	F	M00055775B:G02	UC2-NormColon
4372	509027	1682.M20.gz43_262541	F	M00055778A:G02	UC2-NormColon
4373	546740	1682.F21.gz43_262550	F	M00055767D:F04	UC2-NormColon
4374	559662	1682.L21.gz43_262556	F	M00055775B:G04	UC2-NormColon
4375	388688	1682.B22.gz43_262562	F	M00055764B:F05	UC2-NormColon
4376	555140	1682.G22.gz43_262567	F	M00055768D:G02	UC2-NormColon
4377	640617	1682.K22.gz43_262571	F	M00055774C:D09	UC2-NormColon
4378	447224	1682.D23.gz43_262580	F	M00055766B:B01	UC2-NormColon
4379	644765	1682.B24.gz43_262594	F	M00055764C:C07	UC2-NormColon
4380	648996	1682.F24.gz43_262598	F	M00055768A:B05	UC2-NormColon
4381	642036	1682.I24.gz43_262601	F	M00055772A:C10	UC2-NormColon
4382	644914	1682.J24.gz43_262602	F	M00055773B:A07	UC2-NormColon
4383	458425	1682.M24.gz43_262605	F	M00055778B:E01	UC2-NormColon
4384	556476	1682.N24.gz43_262606	F	M00055779B:F06	UC2-NormColon
4385	672601	1683.E01.gz43_262613	F	M00055785D:C06	UC2-NormColon
4386	463290	1683.H01.gz43_262616	F	M00055789C:F10	UC2-NormColon
4387	675768	1683.D03.gz43_262644	F	M00055784C:H02	UC2-NormColon
4388	642146	1683.H03.gz43_262648	F	M00055789D:B04	UC2-NormColon
4389	648905	1683.F04.gz43_262662	F	M00055787B:F10	UC2-NormColon
4390	467306	1683.I04.gz43_262665	F	M00055790D:G10	UC2-NormColon
4391	497971	1683.C05.gz43_262675	F	M00055783D:B02	UC2-NormColon
4392	645508	1683.N05.gz43_262686	F	M00055795D:E09	UC2-NormColon
4393	644047	1683.005.gz43_262687	F	M00055796B:G05	UC2-NormColon
4394	554117	1683.K06.gz43_262699	F	M00055793A:H09	UC2-NormColon
4395	641919	1683.K08.gz43_262731	F	M00055793B:B06	UC2-NormColon
4396	605761	1683.L08.gz43_262732	F	M00055794A;D08	UC2-NormColon
4397	549434	1683.D09.gz43_262740	F_	M00055785A:H08	UC2-NormColon
4398	446595	1683.M09.gz43_262749	F	M00055795A:F09	UC2-NormColon
4399	643350	1683.F10.gz43_262758	F	M00055787C:D09	UC2-NormColon
4400	454485	1683.M10.gz43_262765	F	M00055795A:F12	UC2-NormColon
4401	647639	1683.O10.gz43_262767	F	M00055796C:E11	UC2-NormColon
4402	510545	1683.I11.gz43_262777	F	M00055791B:E02	UC2-NormColon
4403	77144	1683.N11.gz43_262782	F	M00055795D;H08	UC2-NormColon
4404	140909	1683.C12.gz43_262787	F	M00055784A:B07	UC2-NormColon
4405	650076	1683.D12.gz43_262788	F	M00055785B:B06	UC2-NormColon
4406	449836	1683.F12.gz43_262790	F	M00055787C:E12	UC2-NormColon
4407	642184	1683.B14.gz43_262818	F	M00055783A;C06	UC2-NormColon
4408	638962	1683.H14.gz43_262824	F	M00055790B:A08	UC2-NormColon
4409	641700	1683.M15.gz43_262845	F	M00055795B:F09	UC2-NormColon
4410	437580	1683.N15.gz43_262846	F	M00055796A:A08	UC2-NormColon

Table 2

	able 2				
SEQ ID			ODE		
	CLUSTER	CEO MAME	ORIE NT	CI ONTE ID	T TOD A DAY
		SEQ NAME		CLONE ID	LIBRARY
4411	456626	1683.C16.gz43_262851	F	M00055784A:D05	UC2-NormColon
4412	553979	1683.O16.gz43_262863	F	M00055796D:E06	UC2-NormColon
4413	640525	1683.O17.gz43_262879	F	M00055796D:E10	UC2-NormColon
4414	553546	1683.H18.gz43_262888	F	M00055790B:D05	UC2-NormColon
4415	643230	1683,A19,gz43_262897	F	M00055782B:C08	UC2-NormColon
4416	523590	1683.B20.gz43_262914	F	M00055783B:A04	UC2-NormColon
4417	388085	1683.M20.gz43_262925	F	M00055795C:B10	UC2-NormColon
4418	457396	1683.L21.gz43_262940	F	M00055794C:D10	UC2-NormColon
4419	417617	1683.O23.gz43-262975	F	M00055797B:A11	UC2-NormColon
4420	463821	1684.G01.gz43_262999	F	M00055804D:F02	UC2-NormColon
4421	448787	1684,M01.gz43_263005	F	M00055811C:A01	UC2-NormColon
4422	32812	1684.I03.gz43_263033	F	M00055807A:B10	UC2-NormColon
4423	418340	1684.A04.gz43_263041	F	M00055798B:D12	UC2-NormColon
4424	462249	1684.C04.gz43_263043	F	M00055800D:B03	UC2-NormColon
4425	642099	1684.P04.gz43_263056	F	M00055815C:B03	UC2-NormColon
4426	644915	1684.A05.gz43 263057	F	M00055798B:F02	UC2-NormColon
4427	640222	1684.G05.gz43 263063	F	M00055805A:A02	UC2-NormColon
4428	550370	1684.K05.gz43 263067	F	M00055809A;B10	UC2-NormColon
4429	645560	1684.A06.gz43 263073	F	M00055798B:G04	UC2-NormColon
4430	513238	1684.E06.gz43 263077	F	M00055803A:C06	UC2-NormColon
4431	650231	1684.M07.gz43 263101	F	M00055811D:C02	UC2-NormColon
4432	644314	1684.H08.gz43 263112	F	M00055806B;B10	UC2-NormColon
4433	650782	1684.J08.gz43 263114	F	M00055808B:A04	UC2-NormColon
4434	648221	1684.P08.gz43 263120	F	M00055815C:G05	UC2-NormColon
4435	554742	1684.M09.gz43_263133	F	M00055811D:C12	UC2-NormColon
4436	642962	1684.P09.gz43 263136	F	M00055815D:A11	UC2-NormColon
4437	455028	1684.A10.gz43 263137	F	M00055798D;A10	UC2-NormColon
4438	447015	1684.P10.gz43 263152	F	M00055815D:B02	UC2-NormColon
4439	642653	1684.E11.gz43 263157	F	M00055803A:G08	UC2-NormColon
4440	647336	1684.O11.gz43 263167	F	M00055814C:C07	UC2-NormColon
4441	549640	1684.P11.gz43 263168	F	M00055815D;B03	UC2-NormColon
4442	642198	1684,G12.gz43_263175	F	M00055805B:C08	UC2-NormColon
4443	520320	1684.I12.gz43 263177	F	M00055807B:F05	UC2-NormColon
4444	644147	1684.N12.gz43 263182	F	M00055813A:D10	UC2-NormColon
4445	530774	1684.E13.gz43 263189	F	M00055803B:A11	UC2-NormColon
4446	375814	1684.H14.gz43 263208	F	M00055806C:E09	UC2-NormColon
4447	446789	1684.I14.gz43 263209	F	M00055807B:G10	UC2-NormColon
4448	210839	1684.J14.gz43 263210	F	M00055807B:G10	UC2-NormColon
4449	490414	1684.B15.gz43_263218	F	M00055800A:F02	UC2-NormColon
4450	503491	1684.M15.gz43 263229	F	M00055812A:E01	UC2-NormColon
4451	486134	1684.D17.gz43_263252	F	M00055802B:H03	UC2-NormColon
4452	639441	1684.I17.gz43 263257	F	M00055802B;105	UC2-NormColon
2	055441	1004.117.5243_203237		14100055307C.F05	OCZ-MOINICOION

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T	able 2				
SEQ					
ID			ORIE		
NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
4453	551683	1684.N17.gz43_263262	F	M00055813C;E03	UC2-NormColon
4454	649035	1684.P17.gz43_263264	F	M00055816A:C04	UC2-NormColon
4455	639194	1684.F18.gz43_263270	F	M00055804B:C02	UC2-NormColon
4456	454772	1684.K19.gz43_263291	F	M00055809C:E11	UC2-NormColon
4457	649995	1684.L19.gz43 263292	F	M00055810D:G04	UC2-NormColon
4458	466016	1684.M19.gz43_263293	F	M00055812B:F03	UC2-NormColon
4459	494271	1684.N19.gz43_263294	F	M00055813D:B12	UC2-NormColon
4460	460517	1684.J20.gz43_263306	F	M00055808D:C12	UC2-NormColon
4461	470602	1684.I21.gz43_263321	F	M00055807D:C04	UC2-NormColon
4462	640537	1684.B22.gz43 263330	F	M00055800C:E11	UC2-NormColon
4463	643991	1684.D22.gz43_263332	F	M00055802C:E12	UC2-NormColon
4464	564382	1684.C23.gz43_263347	F	M00055801D:E06	UC2-NormColon
4465	529325	1684.G23.gz43_263351	F	M00055805D:H01	UC2-NormColon
4466	455665	1684.L23.gz43_263356	F	M00055811B:A03	UC2-NormColon
4467	645700	1684.A24.gz43 263361	F	M00055799C:D09	UC2-NormColon
4468	520320	1684.J24.gz43 263370	F	M00055808D:F09	UC2-NormColon
4469	648752	1684.M24.gz43 263373	F	M00055812C:B07	UC2-NormColon
4470	642850	1684.O24.gz43 263375	F	M00055815A:H12	UC2-NormColon
4471	556896	1970.H01.gz43 263397	F	M00054702A:C06	UC2-ColonMetLiver
4472	549238	1970.I01.gz43 263398	F	M00054703D:E07	UC2-ColonMetLiver
4473	551941	1970,M01.gz43 263402	F	M00054710D:A02	UC2-ColonMetLiver
4474	552055	1970.G02.gz43 263412	F	M00054699D:A12	UC2-ColonMetLiver
4475	483141	1970.H02.gz43 263413	F	M00054702A:F03	UC2-ColonMetLiver
4476	561434	1970.I02.gz43 263414	F	M00054703D:F04	UC2-ColonMetLiver
4477	453708	1970,K02.gz43 263416	F	M00054707B:B08	UC2-ColonMetLiver
4478	557935	1970.A03.gz43 263422	F	M00054689C:B11	UC2-ColonMetLiver
4479	509202	1970.H03.gz43 263429	F	M00054702A:H03	UC2-ColonMetLiver
4480	549388	1970.B04.gz43 263439	F	M00054690D:G03	UC2-ColonMetLiver
4481	551243	1970.C05.gz43_263456	F	M00054693A:C09	UC2-ColonMetLiver
4482	549557	1970.N05.gz43_263467	F	M00054712C:C07	UC2-ColonMetLiver
4483	551527	1970.A06.gz43_263470	F	M00054689D:E12	UC2-ColonMetLiver
4484	553128	1970.E06.gz43 263474	F	M00054696B;H11	UC2-ColonMetLiver
4485	549151	1970.K06.gz43_263480	F	M00054707C:D02	UC2-ColonMetLiver
4486	32021	1970.O06.gz43_263484	F	M00054714B:F05	UC2-ColonMetLiver
4487	552598	1970.P07.gz43 263501	F	M00054715D:E11	UC2-ColonMetLiver
4488	552051	1970.D08.gz43_263505	F	M00054695A:A12	UC2-ColonMetLiver
4489	491827	1970.J09.gz43_263527	F	M00054706A:G10	UC2-ColonMetLiver
4490	552625	1970.O09.gz43_263532	F	M00054714C:E01	UC2-ColonMetLiver
4491	552073	1970.M11.gz43 263562	F	M00054711C:A08	UC2-ColonMetLiver
4492	552649	1970.E12.gz43_263570	F	M00054697A:E03	UC2-ColonMetLiver
4493	389201	1970.H12.gz43_263573	F	M00054702D:E07	UC2-ColonMetLiver
4494	552682	1970.I12.gz43 263574	F	M00054704D:F02	UC2-ColonMetLiver

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	able 2				
SEQ					
ID			ORIE		
NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
4495	552686	1970.K12.gz43_263576	F	M00054708A:F11	UC2-ColonMetLiver
4496	559389	1970.A14.gz43_263598	F	M00054690B:B02	UC2-ColonMetLiver
4497	562216	1970.E14.gz43_263602	F	M00054697A:G06	UC2-ColonMetLiver
4498	452392	1970.H14.gz43_263605	F	M00054702D:F05	UC2-ColonMetLiver
4499	552314	1970.I14.gz43_263606	F	M00054705B:C08	UC2-ColonMetLiver
4500	584499	1970.F15.gz43_263619	F	M00054699A:G01	UC2-ColonMetLiver
4501	552418	1970.A16.gz43_263630	F	M00054690B:D10	UC2-ColonMetLiver
4502	551996	1970.F16.gz43_263635	F	M00054699B:A05	UC2-ColonMetLiver
4503	549945	1970.H16.gz43_263637	F	M00054702D:H10	UC2-ColonMetLiver
4504	527679	1970.I16.gz43_263638	F	M00054705B:D02	UC2-ColonMetLiver
4505	391511	1970.K16.gz43_263640	F	M00054708B:F04	UC2-ColonMetLiver
4506	490393	1970.P16.gz43_263645	F	M00054716B:D06	UC2-ColonMetLiver
4507	552197	1970.L17.gz43_263657	F	M00054710B:B10	UC2-ColonMetLiver
4508	550975	1970.O17.gz43_263660	F	M00054715A:G02	UC2-ColonMetLiver
4509	498454	1970.I18.gz43_263670	F	M00054705B:E04	UC2-ColonMetLiver
4510	553938	1970.N18.gz43_263675	F	M00054713C:D07	UC2-ColonMetLiver
4511	448332	1970.C19.gz43_263680	F	M00054693D:A08	UC2-ColonMetLiver
4512	550053	1970.J19.gz43_263687	F	M00054706C:B12	UC2-ColonMetLiver
4513	550330	1970.B20.gz43_263695	F	M00054692B:C06	UC2-ColonMetLiver
4514	450755	1970.C20.gz43_263696	F	M00054693D:C04	UC2-ColonMetLiver
4515	420686	1970.B21.gz43_263711	F	M00054692B:D01	UC2-ColonMetLiver
4516	552581	1970.E21.gz43_263714	F	M00054697C:E11	UC2-ColonMetLiver
4517	551995	1970.F21.gz43_263715	F	M00054699C:A04	UC2-ColonMetLiver
4518	521552	1970.H22.gz43_263733	F	M00054703C:F01	UC2-ColonMetLiver
4519	553358	1970,J22,gz43_263735	F	M00054706D:B03	UC2-ColonMetLiver
4520	558768	1970.L22.gz43_263737	F	M00054710C:A12	UC2-ColonMetLiver
4521	42994	1970.B23.gz43_263743	F	M00054692C:B02	UC2-ColonMetLiver
4522	555255	1970.G23.gz43_263748	F	M00054701D:H05	UC2-ColonMetLiver
4523	451032	1970.O23.gz43_263756	F	M00054715C:D05	UC2-ColonMetLiver
4524	554477	1970.I24.gz43_263766	F	M00054705C:D11	UC2-ColonMetLiver
4525	560317	1970.K24.gz43_263768	F	M00054709A:A10	UC2-ColonMetLiver
4526	554900	1993.J01.gz43_263783	F	M00054863D:H05	UC2-ColonMetLiver
4527	560254	1993.L01.gz43_263785	F	M00054866B:C01	UC2-ColonMetLiver
4528	555740	1993.E02.gz43_263794	F	M00054857C:G09	UC2-ColonMetLiver
4529	551212	1993.F02.gz43_263795	F	M00054858D:C04	UC2-ColonMetLiver
4530	554456	1993.G03.gz43_263812	F	M00054859D:E09	UC2-ColonMetLiver
4531	493303	1993.M03.gz43_263818	F	M00054867B:D02	UC2-ColonMctLiver
4532	522648	1993.P03.gz43_263821	F	M00054870C:C05	UC2-ColonMetLiver
4533	559885	1993.A04.gz43_263822	F	M00054852C:G03	UC2-ColonMetLiver
4534	554257	1993.H04.gz43_263829	F	M00054860D:D01	UC2-ColonMetLiver
4535	554524	1993.B05.gz43_263839	F	M00054854A:H06	UC2-ColonMetLiver
4536	554672	1993,C05,gz43 263840	F	M00054855B:E04	UC2-ColonMetLiver

Table 2

	able 2				
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NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
4537	560714	1993.D05.gz43_263841	F	M00054856D:E11	UC2-ColonMetLiver
4538	557852	1993.G05.gz43_263844	F	M00054859D:G04	UC2-ColonMetLiver
4539	553295	1993.J05.gz43_263847	F	M00054864B:B02	UC2-ColonMetLiver
4540	559027	1993.N06.gz43_263867	F	M00054868C:G11	UC2-ColonMetLiver
4541	548861	1993.B07.gz43_263871	F	M00054854B:C04	UC2-ColonMetLiver
4542	554500	1993.E07.gz43_263874	F	M00054857D:E12	UC2-ColonMetLiver
4543	522507	1993.I07.gz43_263878	F	M00054862C:D05	UC2-ColonMetLiver
4544	501534	1993.J08.gz43_263895	F	M00054864B:D08	UC2-ColonMetLiver
4545	556715	1993.A09.gz43_263902	F	M00054852D:D09	UC2-ColonMetLiver
4546	496897	1993.C09.gz43_263904	F	M00054855C:B06	UC2-ColonMetLiver
4547	554084	1993.D09.gz43_263905	F	M00054857A:B11	UC2-ColonMetLiver
4548	554497	1993.E10.gz43_263922	F	M00054858A:B05	UC2-ColonMetLiver
4549	555968	1993.F10.gz43_263923	F	M00054859A:D10	UC2-ColonMetLiver
4550	43642	1993.K10.gz43_263928	F	M00054865B:H04	UC2-ColonMetLiver
4551	595506	1993.L10.gz43_263929	F	M00054866C:G07	UC2-ColonMetLiver
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4576	593343	1993.D19.gz43_264065	F	M00054857B:D08	UC2-ColonMetLiver
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Table 2

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4657 555057 2018.N22.gz43 264507 F M00055054D:E12 UC2-ColonMetLiver 4658 447815 2018.A23.gz43 264510 F M00055054D:C:G08 UC2-ColonMetLiver 4659 561963 2018.N24.gz43 264539 F M00055055A:B03 UC2-ColonMetLiver 4660 551544 2018.P24.gz43 264541 F M00055057B:B01 UC2-ColonMetLiver 4661 555213 2018.D07.gz43 264641 F M00055043B:B01 UC2-ColonMetLiver	4655	557308	2018.N21.gz43 264491	F	M00055054D:A02	
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4661 555213 2018.D07.gz43_264641 F M00055043B:B01 UC2-ColonMetLiver	4659	561963	2018.N24.gz43 264539	F	M00055055A:B03	UC2-ColonMetLiver
	4660	551544	2018.P24.gz43 264541	F	M00055057B:B01	UC2-ColonMetLiver
4662 555010 2018.J07.gz43_264647 F M00055049C:H12 UC2-ColonMetLiver	4661	555213	2018.D07.gz43_264641	F	M00055043B:B01	UC2-ColonMetLiver
	4662	555010	2018.J07.gz43 264647	F	M00055049C:H12	UC2-ColonMetLiver

Table 2

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4664		2018.A08.gz43_264654	F	M00055039B:G11	UC2-ColonMetLiver
4665	465207	2018.H08.gz43_264661	F	M00055047C:F08	UC2-ColonMetLiver
4666	557834	2018,K08.gz43 264664	F	M00055050D:F04	UC2-ColonMetLiver
4667	557882	2018.L08.gz43_264665	F	M00055051D:G01	UC2-ColonMetLiver
4668	551711	2018.B09.gz43_264671	F	M00055041B:C10	UC2-ColonMetLiver
4669	555524	2018.B10.gz43_264687	F	M00055041B:D11	UC2-ColonMetLiver
4670	446739	2018.J10.gz43_264695	F	M00055049D:D10	UC2-ColonMetLiver
4671	385531	2018.P10.gz43_264701	F	M00055056C:H07	UC2-ColonMetLiver
4672	559052	2018.F11.gz43_264707	F	M00055046B:D02	UC2-ColonMetLiver
4673	553108	2018.M11.gz43_264714	F	M00055053B:A02	UC2-ColonMetLiver
4674	397338	2018.A12.gz43_264718	F	M00055039C:D11	UC2-ColonMetLiver
4675	559699	2018.E12.gz43 264722	F	M00055045B:A04	UC2-ColonMetLiver
4676	553975	2018.O12.gz43 264732	F	M00055055C:E08	UC2-ColonMetLiver
4677	561994	2018.B13.gz43 264735	F	M00055041B:F04	UC2-ColonMetLiver
4678	448098	2018.C13.gz43 264736	F	M00055042B:E05	UC2-ColonMetLiver
4679	558254	2018.D13,gz43 264737	F	M00055043D:D10	UC2-ColonMetLiver
4680	557420	2018.F13.gz43 264739	F	M00055046B;E08	UC2-ColonMetLiver
4681	558007	2018.G13.gz43_264740	F	M00055047A:H04	UC2-ColonMetLiver
4682	557525	2018.M13.gz43 264746	F	M00055053B:C02	UC2-ColonMetLiver
4683	450765	2018.C14.gz43 264752	F	M00055042B:E08	UC2-ColonMetLiver
4684	247	2018.D14.gz43 264753	F	M00055043D:F07	UC2-ColonMetLiver
4685	560720	2018.E14.gz43 264754	F	M00055045B:C08	UC2-ColonMetLiver
4686	491799	2018.F14.gz43 264755	F	M00055046B:F06	UC2-ColonMetLiver
4687	558120	2018.G14.gz43 264756	F	M00055047A:H05	UC2-ColonMetLiver
4688	432970	2018.K14.gz43 264760	F	M00055051A:E11	UC2-ColonMetLiver
4689	561259	2018.A15.gz43 264766	F	M00055039D:D07	UC2-ColonMetLiver
4690	551630	2018.O15.gz43 264780	F	M00055055D:D11	UC2-ColonMetLiver
4691	461	2018.M16.gz43 264794	F	M00055053C:A12	UC2-ColonMetLiver
4692	555399	2018.A17.gz43 264798	F	M00055040A;C02	UC2-ColonMetLiver
4693	554149	2018.B17.gz43 264799	F	M00055041C:C10	UC2-ColonMetLiver
4694	553603	2018.G17.gz43 264804	F	M00055047B:B10	UC2-ColonMetLiver
4695	552641	2018.K17.gz43 264808	F	M00055051B:B08	UC2-ColonMetLiver
4696	562292	2018.M17.gz43 264810	F	M00055053C:B03	UC2-ColonMetLiver
4697	562272	2018.P17.gz43 264813	F	M00055057A;A04	UC2-ColonMetLiver
4698	450755	2018.D18.gz43 264817	F	M00055044A:C02	UC2-ColonMetLiver
4699	555892	2018.E18.gz43 264818	F	M00055045C:F09	UC2-ColonMetLiver
4700	551793	2018.G18.gz43_264820	F	M00055047B:C03	UC2-ColonMetLiver
4701	553002	2018.P18.gz43 264829	·F	M00055057A:A05	UC2-ColonMetLiver
4702	487182	2018.I19.gz43 264838	F	M00055049A:F10	UC2-ColonMetLiver
4703	206098	2018.J19.gz43 264839	F	M00055050B:E11	UC2-ColonMetLiver
4704	557572	2018.K19.gz43 264840	F	M00055051B:D07	UC2-ColonMetLiver

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4706	561046	2018.I20.gz43_264854	F	M00055049A:G03	UC2-ColonMetLiver
4707	454664	2018.L20.gz43_264857	F	M00055052C:B12	UC2-ColonMetLiver
4708	551616	2018.N20.gz43_264859	F	M00055054C:G10	UC2-ColonMetLiver
4709	559294	2018.P20.gz43_264861	F	M00055057A:D10	UC2-ColonMetLiver
4710	548935	2018.C21.gz43_264864	F	M00055042D:G03	UC2-ColonMetLiver
4711	488970	2018.F21.gz43_264867	F	M00055046C:E07	UC2-ColonMetLiver
4712	555751	2018.A22.gz43 264878	F	M00055040B:F02	UC2-ColonMetLiver
4713	423588	2018.C22.gz43 264880	F	M00055042D:H02	UC2-ColonMetLiver
4714	551578	2018.E22.gz43 264882	F	M00055045D:A07	UC2-ColonMetLiver
4715	556171	2018.G22.gz43 264884	F	M00055047B:G06	UC2-ColonMetLiver
4716	451172	2018.P22.gz43 264893	F	M00055057A:F10	UC2-ColonMetLiver
4717	555250	2018.B23.gz43_264895	F	M00055041D:B07	UC2-ColonMetLiver
4718	476268	2018.G23.gz43_264900	F	M00055047B:G10	UC2-ColonMetLiver
4719	553349	2018.K23.gz43 264904	F	M00055051C:B05	UC2-ColonMetLiver
4720	551561	2018.O23.gz43 264908	F	M00055056B:G01	UC2-ColonMetLiver
4721	555173	2018.C24.gz43 264912	F	M00055043A:B06	UC2-ColonMetLiver
4722	557382	2018.L24.gz43 264921	F	M00055052D:B05	UC2-ColonMetLiver
4723	555616	2020.H01.gz43_264933	F	M00055081A:E08	UC2-ColonMetLiver
4724	549607	2020.J01.gz43 264935	F	M00055083B:E05	UC2-ColonMetLiver
4725	492893	2020.L01.gz43 264937	F	M00055086A;B10	UC2-ColonMetLiver
4726	556471	2020.E02.gz43 264946	F	M00055077D:B01	UC2-ColonMetLiver
4727	556542	2020.L02.gz43 264953	F	M00055086A;C09	UC2-ColonMetLiver
4728	551283	2020.N02.gz43 264955	F	M00055088D:A01	UC2-ColonMetLiver
4729	558720	2020.C03.gz43 264960	F	M00055076A:C06	UC2-ColonMetLiver
4730	555512	2020.A04.gz43 264974	F	M00055073D:F06	UC2-ColonMetLiver
4731	549129	2020.D04.gz43 264977	F	M00055077A:B07	UC2-ColonMetLiver
4732	556497	2020,J04,gz43 264983	F	M00055083C:C05	UC2-ColonMetLiver
4733	561422	2020.N04.gz43 264987	F	M00055088D:B08	UC2-ColonMetLiver
4734	556408	2020.O04.gz43 264988	F	M00055090C:B03	UC2-ColonMetLiver
4735	556635	2020.B05.gz43 264991	F	M00055075A:C09	UC2-ColonMetLiver
4736	555368	2020.C05.gz43_264991	F	M00055076A;D11	UC2-ColonMetLiver
4737	495408	2020.C03.gz43_204992 2020.F06.gz43_265011	F	M00055078D:G04	UC2-ColonMetLiver
4738	552361	2020.H06.gz43 265013	F	M00055081B;E10	UC2-ColonMetLiver
4739	556446	2020.K06.gz43 265016	F	M00055084D:B01	
4740	557476	2020.L06.gz43_205017	F	M00055086B:D10	UC2-ColonMetLiver
4741	556559	2020.D06.gz43_265017 2020.O06.gz43_265020	F	M00055086B;D10	UC2-ColonMetLiver UC2-ColonMetLiver
4742	548943	2020.K07.gz43 265032	F		
4742	555126		F	M00055084D:C09	UC2-ColonMetLiver
4744	142614	2020.P07.gz43_265037 2020.E08.gz43_265042	F	M00055091B:A07 M00055078A:C05	UC2-ColonMetLiver
4744	453756		F		UC2-ColonMetLiver
4745	556040	2020.M08.gz43_265050	F	M00055087D:D08	UC2-ColonMetLiver
4/46	330040	2020.G09.gz43_265060	r	M00055080B:G10	UC2-ColonMetLiver

Table 2

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4748	452822	2020.C11.gz43 265088	F	M00055076B;E08	UC2-ColonMetLiver
4749	464905	2020.E11.gz43 265090	F	M00055078A;E10	UC2-ColonMetLiver
4750	509505	2020.F11.gz43_265091	F	M00055079A:H05	UC2-ColonMetLiver
4751	555061	2020.H11.gz43 265093	F	M00055081C:A12	UC2-ColonMetLiver
4752	556668	2020.J11.gz43 265095	F	M00055083D:D08	UC2-ColonMetLiver
4753	549810	2020.C12.gz43_265104	F	M00055076B:F04	UC2-ColonMetLiver
4754	557760	2020.E12.gz43_265106	F	M00055078A:F01	UC2-ColonMetLiver
4755	554084	2020.N12.gz43_265115	F	M00055089B:C01	UC2-ColonMetLiver
4756	16092	2020.P12.gz43_265117	F	M00055091C:B04	UC2-ColonMetLiver
4757	551342	2020.L13.gz43_265129	F	M00055086D:C07	UC2-ColonMetLiver
4758	455820	2020.A14.gz43_265134	F	M00055074B:E05	UC2-ColonMetLiver
4759	555710	2020.I14.gz43_265142	F	M00055082D:E08	UC2-ColonMetLiver
4760	497086	2020.O14.gz43_265148	F	M00055090D:E03	UC2-ColonMetLiver
4761	561489	2020.E15.gz43_265154	F	M00055078B:F05	UC2-ColonMetLiver
4762	558616	2020.F15.gz43_265155	F	M00055079C:G06	UC2-ColonMetLiver
4763	559071	2020.P15.gz43_265165	F	M00055091C:D11	UC2-ColonMetLiver
4764	550701	2020.C16.gz43_265168	F	M00055076B:H06	UC2-ColonMetLiver
4765	551976	2020.G16.gz43_265172	F	M00055080D:A01	UC2-ColonMctLiver
4766	460244	2020.I16.gz43_265174	F	M00055082D:G01	UC2-ColonMetLiver
4767	556802	2020.J16.gz43_265175	F	M00055084A:E10	UC2-ColonMetLiver
4768	454910	2020.K16.gz43_265176	F	M00055085B:D02	UC2-ColonMetLiver
4769	557214	2020.L16.gz43_265177	F	M00055086D:H07	UC2-ColonMetLiver
4770	612961	2020.F17.gz43_265187	F	M00055079D:A03	UC2-ColonMetLiver
4771	465446	2020.I17.gz43_265190	F	M00055082D:H02	UC2-ColonMetLiver
4772	556925	2020.J17.gz43_265191	F	M00055084A:F10	UC2-ColonMetLiver
4773	98869	2020.O17.gz43_265196	F	M00055090D:F03	UC2-ColonMetLiver
4774	553372	2020,H18.gz43_265205	F	M00055081C:G01	UC2-ColonMetLiver
4775	558231	2020.J18.gz43_265207	F	M00055084B:A04	UC2-ColonMetLiver
4 7 76	553318	2020.G19.gz43_265220	F	M00055080D:E07	UC2-ColonMetLiver
4777	449613	2020.H19.gz43_265221	F	M00055081C:H04	UC2-ColonMetLiver
4778	560507	2020.N19.gz43_265227	F	M00055089C:D06	UC2-ColonMetLiver
4779	560069	2020.B20.gz43_265231	F	M00055075D;D05	UC2-ColonMetLiver
4780	557954	2020.C20.gz43_265232	F	M00055076C:H07	UC2-ColonMetLiver
4781	555996	2020.E21.gz43_265250	F	M00055078D:A07	UC2-ColonMetLiver
4782	494625	2020.G21.gz43_265252	F	M00055080D:F01	UC2-ColonMetLiver
4783	554828	2020.P21.gz43_265261	F	M00055091D:A03	UC2-ColonMetLiver
4784	562229	2020.A22.gz43_265262	F	M00055074D:B04	UC2-ColonMetLiver
4785	561975	2020.M22.gz43_265274	F	M00055088C:D01	UC2-ColonMetLiver
4786	560080	2020.N22.gz43_265275	F	M00055090A:F02	UC2-ColonMetLiver
4787	557783	2020.C23.gz43_265280	F	M00055076D:F11	UC2-ColonMetLiver
4788	556881	2020.K23.gz43_265288	F	M00055085D:F03	UC2-ColonMetLiver

Table 2

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4790	598101	2020.023.gz43_265292	F	M00055091A:C09	UC2-ColonMetLiver
4791	451185	2020.F24.gz43_265299	F	M00055080A:F05	UC2-ColonMetLiver
4792	552019	2020.H24.gz43_265301	F	M00055082A:A12	UC2-ColonMetLiver
4793	555725	2020.K24.gz43_265304	F	M00055085D:G09	UC2-ColonMetLiver
4794	562748	2029.P01.gz43_265325	F	M00055111A:F10	UC2-ColonMetLiver
4795	562989	2029.I02.gz43_265334	F	M00055100C:H06	UC2-ColonMetLiver
4796	562243	2029.I03.gz43_265350	F	M00055100D:B02	UC2-ColonMetLiver
4797	562768	2029.B06.gz43_265391	F	M00055093A:F07	UC2-ColonMetLiver
4798	491127	2029.D06.gz43_265393	F	M00055094D:F09	UC2-ColonMetLiver
4799	555330	2029.H07.gz43_265413	F	M00055100A:C05	UC2-ColonMetLiver
4800	552704	2029.L07.gz43_265417	F	M00055105D:B06	UC2-ColonMetLiver
4801	561426	2029.008.gz43_265436	F	M00055110A:C03	UC2-ColonMetLiver
4802	551967	2029.E09.gz43_265442	F	M00055096A:G08	UC2-ColonMetLiver
4803	453533	2029.F09.gz43_265443	F	M00055097B:B12	UC2-ColonMetLiver
4804	394189	2029.009.gz43_265452	F	M00055110A:C05	UC2-ColonMetLiver
4805	562719	2029.F10.gz43_265459	F	M00055097B:F08	UC2-ColonMetLiver
4806	562307	2029.C11.gz43_265472	F	M00055094B;B11	UC2-ColonMetLiver
4807	98484	2029.D11.gz43_265473	F	M00055095A;D08	UC2-ColonMetLiver
4808	558118	2029.G11.gz43_265476	F	M00055099A:G05	UC2-ColonMetLiver
4809	562881	2029.B14.gz43_265519	F	M00055093B:G08	UC2-ColonMetLiver
4810	553548	2029.K14.gz43_265528	F	M00055104B:F09	UC2-ColonMetLiver
4811	637966	2029.A15.gz43_265534	F	M00055092B:G09	UC2-ColonMetLiver
4812	560575	2029.L15.gz43_265545	F	M00055106A:E04	UC2-ColonMetLiver
4813	554496	2029.E16.gz43_265554	F	M00055096C:C03	UC2-ColonMetLiver
4814	561144	2029.J16.gz43_265559	F	M00055103A:H11	UC2-ColonMetLiver
4815	560628	2029.I20.gz43_265622	F	M00055102A:E11	UC2-ColonMetLiver
4816	558511	2029.J20.gz43_265623	F	M00055103C:B07	UC2-ColonMetLiver
4817	402488	2029.C21.gz43_265632	F	M00055094C:C10	UC2-ColonMetLiver
4818	556613	2029.J21.gz43_265639	F	M00055103C:D05	UC2-ColonMetLiver
4819	560898	2029.J22.gz43_265655	F	M00055103C:G03	UC2-ColonMctLiver
4820	454499	2029.L22.gz43_265657	F	M00055106C:B06	UC2-ColonMetLiver
4821	559955	2029.K23.gz43_265672	F	M00055105A:A05	UC2-ColonMetLiver
4822	554233	2030.J01.gz43_265703	F	M00055128A:C10	UC2-ColonMctLiver
4823	557615	2030.K01.gz43_265704	F	M00055128D:D04	UC2-ColonMetLiver
4824	449035	2030.C02.gz43_265712	F	M00055115C:G09	UC2-ColonMetLiver
4825	452525	2030.E03.gz43_265730	F	M00055118C:B03	UC2-ColonMetLiver
4826	452775	2030.L03.gz43_265737	F	M00055129D:C02	UC2-ColonMetLiver
4827	560868	2030.E05.gz43_265762	F	M00055118D:B04	UC2-ColonMctLiver
4828	538830	2030.H05.gz43_265765	F	M00055125A:A02	UC2-ColonMetLiver
4829	486683	2030.K05.gz43_265768	F	M00055129A:B03	UC2-ColonMetLiver
4830	488030	2030.L06.gz43_265785	F	M00055129D:F11	UC2-ColonMetLiver

Table 2

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4832	528369	2030.J07.gz43_265799	F	M00055128B:B12	UC2-ColonMetLiver
4833	479732	2030.G08.gz43_265812	F	M00055124A:F01	UC2-ColonMetLiver
4834	557747	2030.O08.gz43_265820	F	M00055133B:E08	UC2-ColonMetLiver
4835	558534	2030.H09.gz43_265829	F	M00055125B:F01	UC2-ColonMetLiver
4836	312036	2030.D10.gz43_265841	F	M00055117A:G08	UC2-ColonMetLiver
4837	461653	2030.J10.gz43_265847	F	M00055128B:E12	UC2-ColonMetLiver
4838	558060	2030.G11.gz43_265860	F	M00055124B:A11	UC2-ColonMetLiver
4839	557610	2030.L11.gz43_265865	F	M00055130A:D10	UC2-ColonMetLiver
4840	557284	2030.P11.gz43_265869	F	M00055134C:A01	UC2-ColonMetLiver
4841	562714	2030.A12.gz43_265870	F	M00055113B:F02	UC2-ColonMetLiver
4842	453274	2030.C12.gz43_265872	F ·	M00055116A:C07	UC2-ColonMetLiver
4843	551068	2030.P12.gz43_265885	F	M00055134C:B01	UC2-ColonMetLiver
4844	125543	2030.J13.gz43_265895	F	M00055128B:G01	UC2-ColonMetLiver
4845	449454	2030.M13.gz43_265898	F	M00055131A:D07	UC2-ColonMetLiver
4846	562760	2030.O13.gz43_265900	F	M00055133C:C06	UC2-ColonMetLiver
4847	557710	2030.P13.gz43_265901	F	M00055134C:E09	UC2-ColonMetLiver
4848	526334	2030.E14.gz43_265906	F	M00055119D:F08	UC2-ColonMetLiver
4849	451392	2030.O15.gz43_265932	F	M00055133C:G07	UC2-ColonMetLiver
4850	549591	2030.D16.gz43_265937	F	M00055117C:C03	UC2-ColonMetLiver
4851	557895	2030.H16.gz43_265941	F	M00055125C:H03	UC2-ColonMetLiver
4852	554989	2030.J16.gz43_265943	F	M00055128C:E03	UC2-ColonMetLiver
4853	561265	2030.D17.gz43_265953	F	M00055117C:F02	UC2-ColonMetLiver
4854	558015	2030.O17.gz43_265964	F	M00055133C:H11	UC2-ColonMetLiver
4855	561741	2030.A18.gz43_265966	F	M00055114A:E02	UC2-ColonMetLiver
4856	477295	2030.O18.gz43_265980	F	M00055133D:A02	UC2-ColonMetLiver
4857	556310	2030.E19.gz43_265986	F	M00055120B:F12	UC2-ColonMetLiver
4858	558452	2030.H20.gz43_266005	F	M00055125D:E02	UC2-ColonMetLiver
4859	610269	2030.N21.gz43_266027	F	M00055132D:E07	UC2-ColonMetLiver
4860	561279	2030.J22.gz43_266039	F	M00055128D:C11	UC2-ColonMetLiver
4861	557719	2030.N22.gz43_266043	F	M00055132D:E10	UC2-ColonMetLiver
4862	451391	2030.022.gz43_266044	F	M00055133D:F02	UC2-ColonMetLiver
4863	554737	2030.D24.gz43_266065	F	M00055118B:A09	UC2-ColonMetLiver
4864	452759	2030.K24.gz43_266072	F	M00055129C:H08	UC2-ColonMetLiver
4865	562683	2031.A01.gz43_266078	F	M00055135A:E07	UC2-ColonMetLiver
4866	558463	2031.C04.gz43_266128	F	M00055138A:E08	UC2-ColonMetLiver
4867	211273	2031.I04.gz43_266134	F	M00055146A:B12	UC2-ColonMetLiver
4868	468257	2031.N04.gz43_266139	F	M00055152D:C03	UC2-ColonMetLiver
4869	558768	2031.B05.gz43_266143	F	M00055136D:D09	UC2-ColonMetLiver
4870	561245	2031.G05.gz43_266148	F	M00055144A:A11	UC2-ColonMetLiver
4871	557783	2031.K05.gz43_266152	F	M00055149B:F09	UC2-ColonMetLiver
4872	549930	2031.O05.gz43_266156	F	M00055154D:F06	UC2-ColonMetLiver

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4874	450630	2031.G06.gz43_266164	F	M00055144A:E09	UC2-ColonMetLiver
4875	524721	2031.006.gz43_266172	F	M00055154D:G12	UC2-ColonMetLiver
4876	561624	2031.P06.gz43_266173	F	M00055156A:D02	UC2-ColonMetLiver
4877	497493	2031.N07.gz43_266187	F	M00055152D:H09	UC2-ColonMetLiver
4878	490846	2031.M08.gz43_266202	F	M00055152A:B05	UC2-ColonMetLiver
4879	558645	2031.P08.gz43_266205	F	M00055156B:C11	UC2-ColonMetLiver
4880	553969	2031.D09.gz43_266209	F	M00055139B:E10	UC2-ColonMetLiver
4881	558007	2031.P09.gz43_266221	F	M00055156C:A08	UC2-ColonMetLiver
4882	550047	2031.B10.gz43_266223	F	M00055137A:E05	UC2-ColonMetLiver
4883	562000	2031.I10.gz43_266230	F	M00055146B:E09	UC2-ColonMetLiver
4884	160289	2031.L10.gz43_266233	F	M00055150D:C06	UC2-ColonMetLiver
4885	27586	2031.C11.gz43_266240	F	M00055138B:H12	UC2-ColonMetLiver
4886	558230	2031.D11.gz43_266241	F	M00055139B:G03	UC2-ColonMetLiver
4887	549739	2031.M12.gz43_266266	F	M00055152A;E09	UC2-ColonMetLiver
4888	561558	2031.P12.gz43_266269	F	M00055156C:D06	UC2-ColonMctLiver
4889	556630	2031.B14.gz43_266287	F	M00055137B:B11	UC2-ColonMetLiver
4890	561178	2031.P15.gz43_266317	F	M00055156D:A02	UC2-ColonMetLiver
4891	558477	2031.B16.gz43_266319	F	M00055137B:F12	UC2-ColonMetLiver
4892	557250	2031.M16.gz43_266330	F	M00055152B:A03	UC2-ColonMetLiver
4893	556288	2031.P17.gz43_266349	F	M00055157A:B04	UC2-ColonMetLiver
4894	610893	2031.L18.gz43_266361	F	M00055151A:F10	UC2-ColonMetLiver
4895	557708	2031.O24.gz43_266460	F	M00055155D:B02	UC2-ColonMetLiver
4896	562881	2032.E01.gz43_266466	F	M00055162A:B03	UC2-ColonMetLiver
4897	494198	2032.L01.gz43_266473	F	M00055170D:B09	UC2-ColonMetLiver
4898	551475	2032.O01.gz43_266476	F	M00055177A:F05	UC2-ColonMetLiver
4899	558720	2032.K02.gz43_266488	F	M00055169D:A11	UC2-ColonMetLiver
4900	2435	2032.E03.gz43_266498	F	M00055162A:C12	UC2-ColonMetLiver
4901	559464	2032.I04.gz43_266518	F	M00055166D:F02	UC2-ColonMetLiver
4902	559389	2032.A05.gz43_266526	F	M00055157C:C11	UC2-ColonMetLiver
4903	559562	2032.H05.gz43_266533	F	M00055165B:G09	UC2-ColonMetLiver
4904	402799	2032,M05,gz43_266538	F	M00055172A:C09	UC2-ColonMetLiver
4905	557316	2032.J06.gz43_266551	F	M00055168B:F11	UC2-ColonMetLiver
4906	557209	2032.K06.gz43_266552	F	M00055170A:A09	UC2-ColonMetLiver
4907	554604	2032.M06.gz43_266554	F	M00055172A:F03	UC2-ColonMetLiver
4908	558917	2032.H07.gz43_266565	F	M00055165C:H08	UC2-ColonMetLiver
4909	493261	2032.I07.gz43_266566	F	M00055167A;A02	UC2-ColonMetLiver
4910	558755	2032.K07.gz43_266568	F	M00055170A:A11	UC2-ColonMetLiver
4911	558900	2032.M09.gz43_266602	F	M00055172B:B04	UC2-ColonMetLiver
4912	553537	2032.C10.gz43_266608	F	M00055160C:D02	UC2-ColonMetLiver
4913	561413	2032.E10.gz43_266610	F	M00055162B:B04	UC2-ColonMetLiver
4914	561876	2032.F10.gz43_266611	F	M00055163B;F07	UC2-ColonMetLiver

Table 2

	able 2				
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	CLUSTER		NT	CLONE ID	LIBRARY
4915	553797	2032.H10.gz43_266613	F	M00055165D:C03	UC2-ColonMetLiver
4916	558965	2032.G11.gz43_266628	F	M00055164C:C10	UC2-ColonMctLiver
4917	561911	2032.B12.gz43_266639	F	M00055159B:G09	UC2-ColonMetLiver
4918	557961	2032.I12.gz43_266646	F	M00055167B:A08	UC2-ColonMetLiver
4919	558022	2032.K12.gz43_266648	F	M00055170B:B06	UC2-ColonMetLiver
4920	409262	2032.M12.gz43_266650	F	M00055172B:H07	UC2-ColonMetLiver
4921	415538	2032,I15.gz43_266694	F	M00055167B:H07	UC2-ColonMetLiver
4922	559057	2032.M15.gz43_266698	F	M00055172D:D04	UC2-ColonMetLiver
4923	215005	2032.B16.gz43_266703	F	M00055159C:B02	UC2-ColonMetLiver
4924	556542	2032.M16.gz43_266714	F	M00055172D:D07	UC2-ColonMetLiver
4925	562989	2032.O17.gz43_266732	F	M00055177D:F07	UC2-ColonMetLiver
4926	394772	2032.A19.gz43_266750	F	M00055158D:C01	UC2-ColonMetLiver
4927	553877	2032.E19.gz43_266754	F	M00055162C:E12	UC2-ColonMetLiver
4928	558858	2032.I19.gz43_266758	F	M00055167D:B05	UC2-ColonMetLiver
4929	559355	2032.J19.gz43_266759	F	M00055169B:F04	UC2-ColonMetLiver
4930	551693	2032.M19.gz43_266762	F	M00055172D:F12	UC2-ColonMetLiver
4931	323165	2032.E20.gz43_266770	F	M00055162C:G03	UC2-ColonMetLiver
4932	558981	2032.O20.gz43_266780	F	M00055178A:C07	UC2-ColonMetLiver
4933	561507	2032.C21.gz43_266784	F	M00055161A:C02	UC2-ColonMetLiver
4934	555883	2032.E22.gz43_266802	F	M00055162D:B01	UC2-ColonMetLiver
4935	559088	2032.O22.gz43_266812	F	M00055178A:D03	UC2-ColonMetLiver
4936	558395	2032.C23.gz43_266816	F	M00055161A:E05	UC2-ColonMetLiver
4937	447386	2032.D23.gz43_266817	F	M00055161D:H03	UC2-ColonMetLiver
4938	450566	2032.L23.gz43_266825	F	M00055171C:C01	UC2-ColonMetLiver
4939	561918	2032.E24.gz43_266834	F	M00055162D:G04	UC2-ColonMetLiver
4940	559389	2032.G24.gz43_266836	F	M00055165A:F05	UC2-ColonMetLiver
4941	555202	2032.M24.gz43_266842	F	M00055174A:H12	UC2-ColonMetLiver
4942	559752	2032.N24.gz43_266843	F	M00055176D:H01	UC2-ColonMetLiver
4943	478511	2041.B01.gz43_266847	F	M00055181B:A10	UC2-ColonMetLiver
4944	456517	2041.D01.gz43_266849	F	M00055183A:C06	UC2-ColonMetLiver
4945	557895	2041.H01.gz43_266853	F	M00055187D:G11	UC2-ColonMetLiver
4946	561975	2041.A02.gz43_266862	F	M00055179A:G08	UC2-ColonMetLiver
4947	557714	2041.D02.gz43_266865	F	M00055183A:E10	UC2-ColonMetLiver
4948	550874	2041.K02.gz43_266872	F	M00055192C:E04	UC2-ColonMetLiver
4949	539142	2041.L02.gz43_266873	F	M00055193C:C11	UC2-ColonMetLiver
4950	549911	2041.M02.gz43_266874	F	M00055194D:C05	UC2-ColonMetLiver
4951	561830	2041.N02.gz43_266875	F	M00055195C:H05	UC2-ColonMetLiver
4952	479851	2041.A03.gz43_266878	F	M00055179A:H11	UC2-ColonMetLiver
4953	549472	2041.B03.gz43_266879	F	M00055181B:E06	UC2-ColonMetLiver
4954	561687	2041.F03.gz43_266883	F	M00055185D:A02	UC2-ColonMetLiver
4955	504880	2041.L03.gz43_266889	F	M00055193C:E10	UC2-ColonMetLiver
4956	558212	2041.E04.gz43_266898	F	M00055184C:C07	UC2-ColonMetLiver

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	able 2				
SEQ					
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NO	CLUSTER		NT	CLONE ID	LIBRARY
4957	559050	2041.A05.gz43_266910	F	M00055179B:D05	UC2-ColonMetLiver
4958	554520	2041.C05.gz43_266912	F	M00055182A:D07	UC2-ColonMetLiver
4959	409612	2041.E05.gz43_266914	F	M00055184C:D02	UC2-ColonMetLiver
4960	488613	2041.J05.gz43_266919	F	M00055191B:A10	UC2-ColonMetLiver
4961	558326	2041.K05.gz43_266920	F	M00055192C:H06	UC2-ColonMetLiver
4962	555394	2041.F06.gz43_266931	F	M00055185D:D11	UC2-ColonMetLiver
4963	559610	2041.H06.gz43_266933	F	M00055188A:G11	UC2-ColonMetLiver
4964	560080	2041.I06.gz43_266934	F	M00055189B:B12	UC2-ColonMetLiver
4965	555742	2041.N06.gz43_266939	F	M00055195D:B10	UC2-ColonMetLiver
4966	460727	2041.P06.gz43_266941	F	M00055198A:E05	UC2-ColonMetLiver
4967	476199	2041.B07.gz43_266943	F	M00055181C:B07	UC2-ColonMetLiver
4968	516484	2041.G07.gz43_266948	F	M00055187A:F02	UC2-ColonMetLiver
4969	187704	2041.H07.gz43_266949	F	M00055188A:H10	UC2-ColonMetLiver
4970	559000	2041.A08.gz43_266958	F	M00055179B:G07	UC2-ColonMetLiver
4971	553002	2041.B08.gz43_266959	F	M00055181C:B12	UC2-ColonMetLiver
4972	226324	2041.C08.gz43_266960	F	M00055182B:C07	UC2-ColonMetLiver
4973	562840	2041.D08.gz43_266961	F	M00055183C:A02	UC2-ColonMetLiver
4974	561513	2041.E08.gz43_266962	F	M00055184C:F01	UC2-ColonMetLiver
4975	477046	2041.F08.gz43_266963	F	M00055185D:F07	UC2-ColonMetLiver
4976	492627	2041.G08.gz43_266964	F	M00055187A:F06	UC2-ColonMetLiver
4977	559262	2041.H08.gz43_266965	F	M00055188B:E06	UC2-ColonMetLiver
4978	561513	2041.B09.gz43_266975	F	M00055181C:C09	UC2-ColonMetLiver
4979	556632	2041.C09.gz43_266976	F	M00055182B:F05	UC2-ColonMetLiver
4980	552629	2041.G09.gz43_266980	F	M00055187A:G02	UC2-ColonMetLiver
4981	549304	2041.M09.gz43_266986	F	M00055195A:B08	UC2-ColonMetLiver
4982	550652	2041.N09.gz43_266987	F	M00055195D:E11	UC2-ColonMetLiver
4983	490154	2041.P09.gz43_266989	F	M00055198B:H08	UC2-ColonMetLiver
4984	559383	2041.A10.gz43_266990	F	M00055179C:F11	UC2-ColonMetLiver
4985	502343	2041.F10.gz43_266995	F	M00055185D:H01	UC2-ColonMetLiver
4986	413915	2041.I10.gz43_266998	F	M00055189C:C01	UC2-ColonMetLiver
4987	495591	2041.L10.gz43_267001	F	M00055194A:A01	UC2-ColonMetLiver
4988	481231	2041.O10.gz43_267004	F	M00055196D:F07	UC2-ColonMetLiver
4989	559710	2041.A11.gz43_267006	F	M00055179C:H02	UC2-ColonMetLiver
4990	562569	2041.B11.gz43_267007	F	M00055181C:D06	UC2-ColonMetLiver
4991	552629	2041.F11.gz43_267011	F	M00055186A:D04	UC2-ColonMetLiver
4992	560003	2041.K11.gz43_267016	F	M00055193A:A08	UC2-ColonMetLiver
4993	550704	2041.M11.gz43_267018	F	M00055195A:C10	UC2-ColonMetLiver
4994	562569	2041.O11.gz43_267020	F	M00055196D:H02	UC2-ColonMetLiver
4995	561963	2041.D12.gz43_267025	F	M00055183C:D07	UC2-ColonMetLiver
4996	557867	2041.F12.gz43_267027	F	M00055186A:E08	UC2-ColonMetLiver
4997	143218	2041.M12.gz43_267034	F	M00055195A:E07	UC2-ColonMetLiver
4998	495074	2041.N12.gz43_267035	F	M00055196A:C04	UC2-ColonMetLiver

Table 2

Т	able 2				
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5000	558413	.2041.L13.gz43_267049	F	M00055194A:E07	UC2-ColonMetLiver
5001	551977	2041.N13.gz43_267051	F	M00055196A:E01	UC2-ColonMetLiver
5002	558582	2041.E14.gz43_267058	F	M00055185A:G11	UC2-ColonMetLiver
5003	549665	2041.H14.gz43_267061	F	M00055188C:F08	UC2-ColonMetLiver
5004	451544	2041.K14.gz43_267064	F	M00055193A:C06	UC2-ColonMetLiver
5005	450623	2041.M14.gz43_267066	F	M00055195B:B09	UC2-ColonMetLiver
5006	551855	2041.G15.gz43_267076	F	M00055187C:C02	UC2-ColonMetLiver
5007	2284	2041.I15.gz43_267078	F	M00055190A:A05	UC2-ColonMetLiver
5008	559883	2041.J15.gz43_267079	F	M00055192A:A09	UC2-ColonMetLiver
5009	614369	2041.M15.gz43_267082	F	M00055195B:C04	UC2-ColonMetLiver
5010	553237	2041.N15.gz43_267083	F	M00055196A:H07	UC2-ColonMetLiver
5011	559883	2041.E16.gz43 267090	F	M00055185B:B01	UC2-ColonMetLiver
5012	481136	2041.F16.gz43 267091	F	M00055186C:A02	UC2-ColonMetLiver
5013	559885	2041.O16.gz43_267100	F	M00055197B:A10	UC2-ColonMetLiver
5014	551415	2041.P16.gz43 267101	F	M00055198D:A12	UC2-ColonMetLiver
5015	561351	2041.B17.gz43 267103	F	M00055181C:H01	UC2-ColonMetLiver
5016	556881	2041.C17.gz43 267104	F	M00055182C:E09	UC2-ColonMetLiver
5017	549588	2041.G17.gz43 267108	F	M00055187C:E07	UC2-ColonMetLiver
5018	365634	2041.N17.gz43 267115	F	M00055196B:A09	UC2-ColonMetLiver
5019	556343	2041.P17.gz43 267117	F	M00055198D:B08	UC2-ColonMetLiver
5020	624044	2041.K18.gz43 267128	F	M00055193B:A08	UC2-ColonMetLiver
5021	562336	2041.N18.gz43 267131	F	M00055196B:C06	UC2-ColonMetLiver
5022	557935	2041.D19.gz43_267137	F	M00055184A:G02	UC2-ColonMetLiver
5023	557298	2041.G19.gz43 267140	F	M00055187D:A08	UC2-ColonMetLiver
5024	559146	2041.H19.gz43 267141	F	M00055188D:D05	UC2-ColonMetLiver
5025	455814	2041.I19.gz43 267142	F	M00055190A:F11	UC2-ColonMetLiver
5026	549829	2041.A20.gz43_267150	F	M00055181A:E01	UC2-ColonMetLiver
5027	473742	2041.B21.gz43_267167	F	M00055181D:D05	UC2-ColonMetLiver
5028	559333	2041.C21.gz43 267168	F	M00055182D:E06	UC2-ColonMetLiver
5029	559825	2041.H21.gz43 267173	F	M00055188D:H03	UC2-ColonMetLiver
5030	625988	2041.O21.gz43 267180	F	M00055197C:D10	UC2-ColonMetLiver
5031	491260	2041.P21.gz43 267181	F	M00055198D:G03	UC2-ColonMetLiver
5032	557644	2041.A22.gz43 267182	F	M00055181A:G02	UC2-ColonMetLiver
5033	411113	2041.E22.gz43 267186	F	M00055185C:B01	UC2-ColonMetLiver
5034	557961	2041.C23.gz43 267200	F	M00055182D:H08	UC2-ColonMetLiver
5035	559194	2041.A24.gz43_267214	F	M00055181A.H01	UC2-ColonMetLiver
5036	558463	2041.H24.gz43_267221	F	M00055189A:C11	UC2-ColonMetLiver
5037	560859	2041.I24.gz43_267222	F	M00055190C:G08	UC2-ColonMetLiver
5038	456697	2041.N24.gz43 267227	F	M00055196C:B07	UC2-ColonMetLiver
5039	559127	2054.E01.gz43_267234	F	M00055288B:D01	UC2-ColonMetLiver
5040	559460	2054.L01.gz43_267241	F	M00055300C:F11	UC2-ColonMetLiver

Table 2

	able 2				
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NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
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5042	1079	2054,M05.gz43_267306	F	M00055302B:F07	UC2-ColonMetLiver
5043	559113	2054.A07.gz43_267326	F	M00055282A:A01	UC2-ColonMetLiver
5044	500337	2054.J13.gz43_267431	F	M00055297D:C02	UC2-ColonMetLiver
5045	558730	2054.E14.gz43_267442	F	M00055288D:A03	UC2-ColonMetLiver
5046	561779	2054.I14.gz43_267446	F	M00055296A:C05	UC2-ColonMetLiver
5047	397581	2054.H17.gz43_267493	F	M00055294B:C03	UC2-ColonMetLiver
5048	560420	2054.O18.gz43_267516	F	M00055305C:D08	UC2-ColonMetLiver
5049	559728	2054.C19.gz43_267520	F	M00055286A:H08	UC2-ColonMetLiver
5050	559531	2054.H20.gz43_267541	F	M00055294B:G01	UC2-ColonMetLiver
5051	23961	2054.K20.gz43_267544	F	M00055300A:B06	UC2-ColonMetLiver
5052	558446	2054.I21.gz43_267558	F	M00055296C:E08	UC2-ColonMetLiver
5053	562236	2055.K01.gz43_267624	F	M00055323D:A12	UC2-ColonMetLiver
5054	559776	2055.N01.gz43_267627	F	M00055330D:H12	UC2-ColonMetLiver
5055	559574	2055.J04.gz43_267671	F	M00055322C:G11	UC2-ColonMetLiver
5056	558890	2055.I05.gz43_267686	F	M00055321B:B10	UC2-ColonMetLiver
5057	552673	2055.J05.gz43_267687	F	M00055322D:A01	UC2-ColonMetLiver
5058	446900	2055.J07.gz43_267719	F	M00055322D:C12	UC2-ColonMetLiver
5059	559112	2055.C08.gz43_267728	F	M00055312A:D11	UC2-ColonMetLiver
5060	440707	2055.C09.gz43_267744	F	M00055312A:E10	UC2-ColonMetLiver
5061	555359	2055.F12.gz43_267795	F	M00055317C:D04	UC2-ColonMetLiver
5062	472188	2055.P12.gz43_267805	F	M00055335D:A03	UC2-ColonMetLiver
5063	617813	2055.E13.gz43_267810	F	M00055315C:A09	UC2-ColonMetLiver
5064	559027	2055.I13.gz43_267814	F	M00055321D:C12	UC2-ColonMetLiver
5065	448677	2055.K13.gz43_267816	F	M00055324C:H10	UC2-ColonMetLiver
5066	555277	2055.K15.gz43_267848	F	M00055324D:B02	UC2-ColonMetLiver
5067	560538	2055.O16.gz43_267868	F	M00055334C:E11	UC2-ColonMetLiver
5068	560369	2055.D17.gz43_267873	F	M00055313D:E10	UC2-ColonMetLiver
5069	559794	2055.G19.gz43_267908	F	M00055319B:H06	UC2-ColonMetLiver
5070	138470	2055.B20.gz43_267919	F	M00055310B;E02	UC2-ColonMetLiver
5071	558105	2055,H20.gz43_267925	F	M00055320D:E09	UC2-ColonMetLiver
5072	473343	2055.K21.gz43_267944	F	M00055325A:E12	UC2-ColonMetLiver
5073	554176	2055.O21.gz43_267948	F	M00055334D:G07	UC2-ColonMetLiver
5074	477757	2055.E22.gz43_267954	F	M00055316B:B10	UC2-ColonMetLiver
5075	553709	2055.G22.gz43_267956	F	M00055319C:C03	UC2-ColonMetLiver
5076	554585	2055.K22.gz43_267960	F	M00055325A:H02	UC2-ColonMetLiver
5077	451429	2055.N22.gz43 267963	F	M00055333C:F12	UC2-ColonMetLiver
5078	551811	2055.A23.gz43_267966	F	M00055308D:C09	UC2-ColonMetLiver
5079	62458	2055.C23.gz43_267968	F	M00055312D:A09	UC2-ColonMetLiver
5080	551912	2055,D23.gz43_267969	F	M00055314B:G07	UC2-ColonMetLiver
5081	552857	2055.G23.gz43_267972	F	M00055319C:C07	UC2-ColonMetLiver
5082	555349	2055.N23.gz43_267979	F	M00055333C:H07	UC2-ColonMetLiver

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SEQ					
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5084	550063	2056.G01.gz43_268004	F	M00055344D:A09	UC2-ColonMetLiver
5085	560544	2056.M01.gz43_268010	F	M00055352B:E06	UC2-ColonMetLiver
5086	561108	2056.P01.gz43_268013	F	M00055356C;H02	UC2-ColonMetLiver
5087	512014	2056.L02.gz43_268025	F	M00055351B:H12	UC2-ColonMetLiver
5088	627386	2056.D03.gz43_268033	F	M00055341C:F04	UC2-ColonMetLiver
5089	561124	2056,E03.gz43_268034	F	M00055342C:H06	UC2-ColonMetLiver
5090	418482	2056.H03.gz43_268037	F	M00055346A:E12	UC2-ColonMetLiver
5091	558562	2056.A04.gz43_268046	F	M00055336D:B03	UC2-ColonMetLiver
5092	551912	2056.B04.gz43_268047	F	M00055338C:F03	UC2-ColonMetLiver
5093	552669	2056.D04.gz43_268049	F	M00055341C:G12	UC2-ColonMetLiver
5094	559857	2056.J04.gz43_268055	F	M00055348D:A01	UC2-ColonMetLiver
5095	557372	2056.M04.gz43_268058	F	M00055352B:H05	UC2-ColonMetLiver
5096	558182	2056.N04.gz43_268059	F	M00055354A:A01	UC2-ColonMetLiver
5097	552430	2056.B05.gz43_268063	F	M00055338C:G04	UC2-ColonMetLiver
5098	560932	2056.F05.gz43_268067	F	M00055343D:G03	UC2-ColonMetLiver
5099	559049	2056.G05.gz43_268068	F	M00055345A:D05	UC2-ColonMetLiver
5100	449405	2056.C06.gz43_268080	F	M00055340B;C03	UC2-ColonMetLiver
5101	558769	2056.G06.gz43_268084	F	M00055345B:B03	UC2-ColonMetLiver
5102	556216	2056.K06.gz43_268088	F	M00055350A:F01	UC2-ColonMetLiver
5103	553591	2056.M06.gz43_268090	F	M00055352C:A07	UC2-ColonMetLiver
5104	561024	2056.B07.gz43_268095	F	M00055338C:H06	UC2-ColonMetLiver
5105	560652	2056.C07.gz43_268096	F	M00055340B:E06	UC2-ColonMetLiver
5106	559296	2056.F07.gz43_268099	F	M00055343D:H04	UC2-ColonMetLiver
5107	560174	2056.P07.gz43_268109	F	M00055356D:C11	UC2-ColonMetLiver
5108	552357	2056.D08.gz43_268113	F	M00055341D:H07	UC2-ColonMetLiver
5109	627139	2056.E08.gz43_268114	·F	M00055342D:F07	UC2-ColonMetLiver
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5200 453846 2065.011.gz43 268556 F M00055376B:B01 UC2-ColonMetLiver 5201 556430 2065.G12.gz43 268564 F M00055367A:B11 UC2-ColonMetLiver 5202 549984 2065.J13.gz43 268567 F M00055370D:P06 UC2-ColonMetLiver 5203 561825 2065.F13.gz43 268579 F M00055376C:F11 UC2-ColonMetLiver 5204 560959 2065.J13.gz43 268583 F M0005537D:H07 UC2-ColonMetLiver 5206 562801 2065.P13.gz43 268586 F M00055377D:H2 UC2-ColonMetLiver 5207 559675 2065.I14.gz43 268593 F M0005537C:G08 UC2-ColonMetLiver	5199			F		
5201 556430 2065.GI2.gz43 268564 F M00055367A:B11 UC2-ColonMetLiver 5202 549984 2055.JI2.gz43 268567 F M00055370D:F06 UC2-ColonMetLiver 5203 561825 2065.F13.gz43 268579 F M00055370D:H07 UC2-ColonMetLiver 5204 560959 2065.J13.gz43 268583 F M00055370D:H07 UC2-ColonMetLiver 5206 56280 2065.P13.gz43 268586 F M00055377D:F12 UC2-ColonMetLiver 5207 559675 2065.D14.gz43 268589 F M00055362:C:G08 UC2-ColonMetLiver						
5202 549984 2065 J12 ge43 268567 F M00055370D:F06 UC2-ColonMetLiver 5203 561825 2065 F13 ge43 268579 F M00055370D:F07 UC2-ColonMetLiver 5204 560959 2065 J13 ge43 268583 F M00055370D:H07 UC2-ColonMetLiver 5205 636876 2065 M13 ge43 268586 F M00055374X:E01 UC2-ColonMetLiver 5206 562801 2065 D14 ge43 268589 F M0005537C:G08 UC2-ColonMetLiver 5207 599675 2065 D14 ge43 268593 F M0005537C:G08 UC2-ColonMetLiver						
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5204 560959 2065.J13.gz43 268583 F M00055370D:H07 UC2-ColonMetLiver 5205 636876 2065.M13.gz43 268586 F M00055374A:E01 UC2-ColonMetLiver 5206 562801 2065.P13.gz43 268589 F M0005537D:F12 UC2-ColonMetLiver 5207 559675 2065.D14.gz43 268593 F M00053362C:G08 UC2-ColonMetLiver						
5205 636876 2065.M13.gz43_26886 F M00055374A:E01 UC2-ColonMetLiver 5206 562801 2065.P13.gz43_268589 F M00055377D:F12 UC2-ColonMetLiver 5207 559675 2065.D14.gz43_268593 F M00055362C:G08 UC2-ColonMetLiver	$\overline{}$					
5206 562801 2065.P13.gz43 268589 F M00055377D:F12 UC2-ColonMetLiver 5207 559675 2065.D14.gz43 268593 F M00055362C:G08 UC2-ColonMetLiver						
5207 559675 2065.D14.gz43 268593 F M00055362C:G08 UC2-ColonMetLiver						
	5208	632260	2065.K14,gz43 268600	F	M00055372B:E01	UC2-ColonMetLiver

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Table 2

T	able 2				
SEQ					
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5209	490308	2065.L14.gz43_268601	F	M00055373B:A09	UC2-ColonMetLiver
5210	559828	2065.B15.gz43_268607	F	M00055359D:H02	UC2-ColonMetLiver
5211	559022	2065.C15.gz43_268608	F	M00055361A:C01	UC2-ColonMetLiver
5212	556850	2065.F15.gz43_268611	F	M00055366A:B04	UC2-ColonMetLiver
5213	508515	2065.J15.gz43_268615	F	M00055371A:B05	UC2-ColonMetLiver
5214	465610	2065.A17.gz43_268638	F	M00055358D:G04	UC2-ColonMetLiver
5215	463304	2065.E17.gz43_268642	F	M00055364B:D01	UC2-ColonMetLiver
5216	551551	2065.F17.gz43_268643	F	M00055366A;H08	UC2-ColonMetLiver
5217	559963	2065.E18.gz43_268658	F	M00055364B:E10	UC2-ColonMetLiver
5218	562027	2065.J18.gz43_268663	F	M00055371A:H10	UC2-ColonMetLiver
5219	561868	2065,K18,gz43_268664	F	M00055372B:F11	UC2-ColonMetLiver
5220	561610	2065.J19.gz43_268679	F	M00055371B:D01	UC2-ColonMetLiver
5221	562565	2065.M19.gz43_268682	F	M00055374B:D05	UC2-ColonMetLiver
5222	235456	2065.P19.gz43_268685	F	M00055378A:B12	UC2-ColonMetLiver
5223	553615	2065.C20.gz43_268688	F	M00055361B:F12	UC2-ColonMetLiver
5224	534054	2065.E20.gz43_268690	F	M00055364C:B08	UC2-ColonMetLiver
5225	511746	2065.F20.gz43 268691	F	M00055366B:C04	UC2-ColonMetLiver
5226	559495	2065.A21.gz43 268702	F	M00055359B;F03	UC2-ColonMetLiver
5227	559675	2065.C21.gz43 268704	F	M00055361B;G08	UC2-ColonMetLiver
5228	553705	2065.J21.gz43 268711	F	M00055371B:F01	UC2-ColonMetLiver
5229	559764	2065.M21.gz43 268714	F	M00055374B:F06	UC2-ColonMetLiver
5230	482090	2065.N21.gz43 268715	F	M00055375B:H02	UC2-ColonMetLiver
5231	562459	2065.C22.gz43 268720	F	M00055361C:E05	UC2-ColonMetLiver
5232	553904	2065.K22.gz43 268728	F	M00055372C:E03	UC2-ColonMetLiver
5233	555878	2065.N22.gz43 268731	F	M00055375B:H07	UC2-ColonMetLiver
5234	559676	2065.A23.gz43 268734	F	M00055359B:G09	UC2-ColonMetLiver
5235	505971	2065.F23.gz43 268739	F	M00055366C:B11	UC2-ColonMetLiver
5236	62458	2065.H23.gz43 268741	F	M00055368D:E03	UC2-ColonMetLiver
5237	207099	2065.O23.gz43 268748	F	M00055377A;B11	UC2-ColonMetLiver
5238	491635	2065.A24.gz43 268750	F	M00055359B:H07	UC2-ColonMetLiver
5239	32021	2065,D24,gz43 268753	F	M00055363A:D02	UC2-ColonMetLiver
5240	448741	2065.N24.gz43 268763	F	M00055375C:C08	UC2-ColonMetLiver
5241	504560	2078.H01.gz43_269031	F	M00042953B:D02	UC3-NormColon
5242	516415	2078.M01.gz43 269036	F	M00042965A:G02	UC3-NormColon
5243	452981	2078.J02.gz43 269049	F	M00042958B:H04	UC3-NormColon
5244	514594	2078.L02.gz43_269051	F	M00042963B:E12	UC3-NormColon
5245	451923	2078.B03.gz43 269057	F	M00042586C:E01	UC3-NormColon
5246	512721	2078.J03.gz43 269065	F	M00042958C:D04	UC3-NormColon
5247	512051	2078.K03.gz43 · 269066	F	M00042960D:C11	UC3-NormColon
5248	506920	2078.P03.gz43 269071	F	M00042976B:F05	UC3-NormColon
5249	451401	2078.F04.gz43 269077	F	M00042628B:G11	UC3-NormColon
5250	448594	2078.I04.gz43 269080	F	M00042955D:H03	UC3-NormColon

Table 2

	able 2				
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5251	452898	2078.M04.gz43 269084			LIBRARY
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		2078.N04.gz43_269085		M00042969B:E01	UC3-NormColon
5253		2078.O04.gz43_269086	F	M00042972C:F07	UC3-NormColon
5254		2078.G05.gz43_269094	F	M00042951D:C05	UC3-NormColon
5255	448090	2078.I05.gz43_269096	F	M00042956A:H02	UC3-NormColon
5256		2078.N05.gz43_269101	F	M00042969B:G10	UC3-NormColon
5257	373239	2078.O05.gz43_269102	F	M00042972C:F09	UC3-NormColon
5258	451802	2078.D06.gz43_269107	F	M00042623B:B09	UC3-NormColon
5259	511746	2078.E06.gz43_269108	F	M00042625D:C07	UC3-NormColon
5260	516522	2078.F06.gz43_269109	F	M00042628C:G10	UC3-NormColon
5261	423578	2078.J06.gz43_269113	F	M00042959A:B07	UC3-NormColon
5262	513888	2078.E07.gz43_269124	F	M00042625D:E08	UC3-NormColon
5263	452801	2078.L07.gz43_269131	F	M00042963D:F11	UC3-NormColon
5264	452775	2078.M07.gz43_269132	F	M00042965C:E06	UC3-NormColon
5265	415825	2078.C08.gz43_269138	F	M00042589D:D08	UC3-NormColon
5266	447904	2078.F08.gz43_269141	F	M00042628D:F12	UC3-NormColon
5267	514142	2078.J08.gz43_269145	F	M00042959A:E08	UC3-NormColon
5268	92639	2078.B09.gz43_269153	F	M00042587B:G07	UC3-NormColon
5269	452182	2078.C09.gz43_269154	F	M00042589D:F02	UC3-NormColon
5270	447210	2078.D09.gz43_269155	F	M00042623D:C02	UC3-NormColon
5271	452500	2078.O09.gz43_269166	F	M00042973B:B10	UC3-NormColon
5272	452204	2078.A10.gz43_269168	F	M00042583D:F03	UC3-NormColon
5273	447025	2078.E10.gz43 269172	F	M00042626B:A12	UC3-NormColon
5274	452324	2078.G10.gz43 269174	F	M00042952A:H12	UC3-NormColon
5275	506901	2078.A11.gz43_269184	F	M00042583D:F11	UC3-NormColon
5276	507349	2078.C11.gz43 269186	F	M00042590B:G02	UC3-NormColon
5277	452052	2078.F11.gz43_269189	F	M00042629A:E11	UC3-NormColon
5278	500853	2078.G11.gz43 269190	F	M00042952B:A08	UC3-NormColon
5279	514160	2078.J11.gz43 269193	F	M00042959B:E11	UC3-NormColon
5280	418340	2078.C12.gz43 269202	F	M00042590C:C09	UC3-NormColon
5281	452142	2078.H12.gz43_269207	F	M00042954A:F04	UC3-NormColon
5282	451994	2078.I12.gz43_269208	F	M00042957A:D06	UC3-NormColon
5283	452615	2078.J12.gz43_269209	F	M00042959C:C06	· UC3-NormColon
5284	452899	2078.K12.gz43 269210	F	M00042962A:G04	UC3-NormColon
5285	446438	2078.K13.gz43 269226	F	M00042962B:A03	UC3-NormColon
5286	452462	2078.D14.gz43 269235	F	M00042624D:B05	UC3-NormColon
5287	510169	2078.J14.gz43 269241	F	M00042959D:A05	UC3-NormColon
5288	508088	2078.B16.gz43 269265	F	M00042588A:G10	UC3-NormColon
5289	452031	2078.E16.gz43 269268	F	M00042626D:D12	UC3-NormColon
5290	513155	2078.K16.gz43 269274	F	M00042962C:D05	UC3-NormColon
5291	513156	2078.L16.gz43 269275	F	M00042964C:D06	UC3-NormColon
5292	452695	2078.N16.gz43 269277	F	M00042970D:D02	UC3-NormColon
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Table 2

ıble 2				
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			M00042964C:D10	UC3-NormColon
510717		F	M00042971A:B09	UC3-NormColon
			M00042974D:B07	UC3-NormColon
		F	M00042978B:F03	UC3-NormColon
		F	M00042627B:A10	UC3-NormColon
			M00042954D:D04	UC3-NormColon
451929	2078.I18.gz43_269304	F	M00042957D:C09	UC3-NormColon
511351	2078.K18.gz43_269306	F	M00042962D:B09	UC3-NormColon
505858	2078.B19.gz43_269313	F	M00042588C:E07	UC3-NormColon
501401	2078.C19.gz43_269314	F	M00042621D:A03	UC3-NormColon
452662	2078.N19.gz43_269325		M00042971A:D12	UC3-NormColon
512059	2078.L21.gz43_269355	F	M00042964D:C02	UC3-NormColon
450805	2078.O21.gz43_269358	F	M00042975C:A08	UC3-NormColon
504007	2078.B22.gz43_269361	F	M00042589B:D04	UC3-NormColon
450262	2078.A23.gz43_269376	F	M00042586B:A09	UC3-NormColon
452066	2078.B23.gz43_269377	F	M00042589B:E03	UC3-NormColon
452459	2078.K23.gz43_269386	F	M00042963B:A02	UC3-NormColon
452506	2078.L23.gz43_269387	F	M00042965A:B03	UC3-NormColon
448453	2078.A24.gz43_269392	F	M00042586B:A10	UC3-NormColon
7022	2089.A01.gz43_269708	F	M00042611A:A01	UC3-PrimColon
523182	2089.E01.gz43_269712	F	M00043079D:G10	UC3-PrimColon
2930	2089.L01.gz43_269719	F	M00043108B:D12	UC3-PrimColon
454701	2089.H02.gz43_269731	F	M00043093D:F12	UC3-PrimColon
454226	2089.I02.gz43_269732	F	M00043097D:B12	UC3-PrimColon
524624	2089.O02.gz43_269738	F	M00043139D:A06	UC3-PrimColon
453220	2089.B03.gz43_269741	F	M00042614D:A12	UC3-PrimColon
453893	2089.C03.gz43_269742	F	M00042618B:G04	UC3-PrimColon
453470	2089.F03.gz43_269745	F	M00043086A:C02	UC3-PrimColon
524478	2089.G03.gz43_269746	F	M00043090D:H07	UC3-PrimColon
529219	2089.H03.gz43_269747	F	M00043094A:F01	UC3-PrimColon
454050	2089.P03.gz43_269755	F	M00043144D:H01	UC3-PrimColon
519988	2089.E04.gz43_269760	F	M00043080B:C11	UC3-PrimColon
523590	2089.J04.gz43_269765	F	M00043102A:G12	UC3-PrimColon
452936	2089.N04.gz43_269769	F	M00043137C:D02	UC3-PrimColon
453761	2089.J05.gz43_269781	F	M00043102B:F05	UC3-PrimColon
405102	2089.A06.gz43_269788	F	M00042612A:B06	UC3-PrimColon
454720	2089.L06.gz43_269799	F	M00043109C:F04	UC3-PrimColon
454438	2089.N06.gz43_269801	F	M00043137D:D10	UC3-PrimColon
448985	2089.D07.gz43_269807	F	M00043077B:H01	UC3-PrimColon
530656	2089.G07.gz43_269810	F	M00043091C:H05	UC3-PrimColon
454134	2089.L07.gz43_269815	F	M00043131B:A11	UC3-PrimColon
	456545 42444 451993 451993 451929 505858 501401 452662 452662 452663 452663 452664 452666 452469 452666 452469 452666 452469 452666 452469 452666 452469 452666 452469 452666 452469 452666 452469 452666 452469 452666 452469 452666 452469 452666 452469 452666 452469 452666 452469 452666 452666 452666 452666 452666 452666 452666 452666 452666 452666 452666 452666 452666 452666 4526666 4526666 4526666 4526666 4526666 4526666 4526666 4526666 45266666 45266666 4526666666666	448332 2078.I17.gz43 269289 513168 2078.I17.gz43 269291 513168 2078.I17.gz43 269294 4244 2078.I17.gz43 269294 4244 2078.I18.gz43 269303 452376 2078.I18.gz43 269303 451993 2078.H18.gz43 269303 451993 2078.H18.gz43 269313 501401 2078.S.R18.gz43 269313 501401 2078.C19.gz43 269313 501401 2078.L19.gz43 269315 452662 2078.N19.gz43 269315 50805 2078.C12.gz43 269355 504007 2078.E22.gz43 269346 452662 2078.R23.gz43 269364 452062 2078.R23.gz43 269346 452062 2078.R23.gz43 269346 452062 2078.R23.gz43 269346 452060 2078.E23.gz43 269347 452566 2078.R23.gz43 269347 452566 2078.R23.gz43 269347	448332 2078.J17.g243 269289 F 513168 2078.L17.g243 269291 F 513168 2078.N17.g243 269294 F 456545 2078.017.g243 269294 F 452376 2078.R18.g243 269300 F 451993 2078.H18.g243 269300 F 451993 2078.H18.g243 269303 F 451993 2078.H18.g243 269304 F 51351 2078.K18.g243 269304 F 50401 2078.C19.g243 269313 F 505858 2078.R19.g243 269315 F 50400 2078.R19.g243 269355 F 504005 2078.C19.g243 269355 F 504005 2078.R21.g243 269361 F 452062 2078.R22.g243 269361 F 452062 2078.R23.g243 269376 F 452066 2078.R23.g243 269376 F 452066 2078.R23.g243	CLUSTER SEQ NAME NT CLONE ID 448332 2078.117.gz43 269289 F M00042996A:B10 510717 2078.N17.gz43 269291 F M0004297La:B06 510717 2078.N17.gz43 269293 F M0004297La:B06 456545 2078.017.gz43 269293 F M0004297B-B07 452376 2078.E18.gz43 269293 F M00042297B-D10 451929 2078.H18.gz43 269300 F M00042295D-D04 451929 2078.B18.gz43 269304 F M00042957D-C09 501451 2078.K18.gz43 269304 F M0004295D-D09 501451 2078.K18.gz43 269304 F M00042957D-C09 501451 2078.K18.gz43 269304 F M0004295D-D09 504804 2078.B19.gz43 269314 F M00042258C-D09 504040 2078.R19.gz43 269314 F M00042258C-D09 512652 2078.N2.l.gz43 269358 F M0004259A-D01 512659 2078.L21.gz43 269358 F M0004258B-D04 450007 2078.B22.gz43 269361 F

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Table 2

	able 2				
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NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
5335	454910	2089.G08.gz43_269826	F	M00043091C:H08	UC3-PrimColon
5336	417549	2089.D09.gz43_269839	F	M00043077D:E12	UC3-PrimColon
5337	454374	2089.G09.gz43_269842	F	M00043091D:C01	UC3-PrimColon
5338	454518	2089.M09.gz43_269848	F	M00043134C:D06	UC3-PrimColon
5339	523753	2089.N09.gz43_269849	F	M00043138A:H03	UC3-PrimColon
5340	522648	2089.D10.gz43_269855	F	M00043077D:F04	UC3-PrimColon
5341	407275	2089.N10.gz43_269865	F	M00043138B:B08	UC3-PrimColon
5342	415326	2089.O10.gz43_269866	F	M00043141C:C12	UC3-PrimColon
5343	453756	2089.P10.gz43_269867	F	M00043146C:F10	UC3-PrimColon
5344	453766	2089.J11.gz43_269877	F	M00043102D:F11	UC3-PrimColon
5345	450287	2089.K11.gz43_269878	F	M00043106B:F07	UC3-PrimColon
5346	454825	2089.L11.gz43_269879	F	M00043131B:G10	UC3-PrimColon
5347	449335	2089.J12.gz43_269893	F	M00043103A:G05	UC3-PrimColon
5348	453572	2089.K12.gz43_269894	F	M00043106C:D05	UC3-PrimColon
5349	454550	2089,M12.gz43_269896	F	M00043135C:E07	UC3-PrimColon
5350	23649	2089.N12.gz43_269897	F	M00043138B:F02	UC3-PrimColon
5351	449430	2089.F13.gz43_269905	F	M00043087B;D10	UC3-PrimColon
5352	528134	2089.H13.gz43_269907	F	M00043096A:E01	UC3-PrimColon
5353	451391	2089.L13.gz43_269911	F	M00043131C:A11	UC3-PrimColon
5354	454126	2089.M13.gz43_269912	F	M00043135D:A11	UC3-PrimColon
5355	437064	2089.N13.gz43_269913	F	M00043138B:G11	UC3-PrimColon
5356	454129	2089.O13.gz43_269914	F	M00043141D:A12	UC3-PrimColon
5357	519109	2089.C14.gz43_269918	F	M00043075A:B12	UC3-PrimColon
5358	519641	2089.D14.gz43_269919	F	M00043078C:C04	UC3-PrimColon
5359	526575	2089.M14.gz43_269928	F	M00043135D:C07	UC3-PrimColon
5360	453783	2089.A15.gz43_269932	F	M00042613A:H01	UC3-PrimColon
5361	453533	2089.I15.gz43_269940	F	M00043100C:D08	UC3-PrimColon
5362	449394	2089.N15.gz43_269945	F	M00043138C:D09	UC3-PrimColon
5363	523732	2089.D16.gz43_269951	F	M00043078C:H05	UC3-PrimColon
5364	395536	2089.F16.gz43_269953	F	M00043088B:D07	UC3-PrimColon
5365	454509	2089.H16.gz43_269955	F	M00043096C:D02	UC3-PrimColon
5366	454562	2089.E17.gz43_269968	F	M00043082D;B05	UC3-PrimColon
5367	451972	2089.H17.gz43_269971	F	M00043096C:H04	UC3-PrimColon
5368	453726	2089.K17.gz43 269974	F	M00043107A;E07	UC3-PrimColon
5369	527361	2089.M17.gz43_269976	F	M00043136A;D03	UC3-PrimColon
5370	453783	2089.A18.gz43_269980	F	M00042613C:F06	UC3-PrimColon
5371	453385	2089.B20.gz43_270013	F	M00042617C:B07	UC3-PrimColon
5372	454177	2089.G20.gz43_270018	F	M00043093A;A06	UC3-PrimColon
5373	801	2089.I20.gz43_270020	F	M00043101A:F12	UC3-PrimColon
5374	453494	2089.J20.gz43_270021	F	M00043104B:C09	UC3-PrimColon
5375	453202	2089.P20.gz43_270027	F	M00043149B:A01	UC3-PrimColon
5376	520616	2089.F21.gz43_270033	F	M00043089A:D06	UC3-PrimColon
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Т	able 2				
SEQ					
ID			ORIE		
NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
5377	527873	2089.H21.gz43_270035	F	M00043097A:D11	UC3-PrimColon
5378	450566	2089.K21.gz43_270038	F	M00043107D:H04	UC3-PrimColon
537.9	520074	2089.A22.gz43_270044	F	M00042614A:C04	UC3-PrimColon
5380	518566	2089.C22.gz43_270046	F	M00043076A:A09	UC3-PrimColon
5381	387530	2089.H22.gz43_270051	F	M00043097A:F06	UC3-PrimColon
5382	453846	2089.I22.gz43_270052	F	M00043101C:F12	UC3-PrimColon
5383	454483	2089.L22.gz43_270055	F	M00043132C:D02	UC3-PrimColon
5384	524157	2089,A23.gz43 270060	F	M00042614A:H02	UC3-PrimColon
5385	453132	2089.K23.gz43 270070	F	M00043108B:A01	UC3-PrimColon
5386	524622	2089.N23.gz43 270073	F	M00043139C:A05	UC3-PrimColon
5387	447446	2089,A24.gz43 270076	F	M00042614C:B09	UC3-PrimColon
5388	524470	2089.F24.gz43 270081	F	M00043090B:H06	UC3-PrimColon
5389	446242	2066.N03.gz43 270439	F	M00055392A:H06	UC2-ColonMetLiver
5390	554908	2066,P07.gz43 270505	F	M00055394D:F03	UC2-ColonMetLiver
5391	551995	2066.F08.gz43 270511	F	M00055383B:H09	UC2-ColonMetLiver
5392	561325	2066.H08.gz43 270513	F	M00055385C:F06	UC2-ColonMetLiver
5393	553244	2066.J08.gz43_270515	F	M00055388A:A09	UC2-ColonMetLiver
5394	562876	2066.D09.gz43 270525	F	M00055381C:G03	UC2-ColonMetLiver
5395	562216	2066.F09.gz43 270527	F	M00055383C:A08	UC2-ColonMetLiver
5396	560948	2066.J09.gz43 270531	F	M00055388A:B06	UC2-ColonMetLiver
5397	446531	2066.P09.gz43 270537	F	M00055395A:C02	UC2-ColonMetLiver
5398	556793	2066.K10.gz43 270548	F	M00055388D:F11	UC2-ColonMetLiver
5399	499690	2066.M10.gz43 270550	F	M00055391B:D05	UC2-ColonMetLiver
5400	558297	2066.I11.gz43_270562	F	M00055386D:G02	UC2-ColonMetLiver
5401	453508	2066.M11.gz43 270566	F	M00055391B:D07	UC2-ColonMetLiver
5402	551640	2066.N11.gz43 270567	F	M00055392C:G07	UC2-ColonMetLiver
5403	562085	2066.C12.gz43 270572	F	M00055380D:H02	UC2-ColonMetLiver
5404	446399	2066.E12.gz43 270574	F	M00055382C:H06	UC2-ColonMetLiver
5405	562498	2066.G12.gz43 270576	F	M00055384B:D10	UC2-ColonMetLiver
5406	562524	2066.A13.gz43 270586	F	M00055378D:D04	UC2-ColonMetLiver
5407	465836	2066.J13.gz43 270595	F	M00055388B;B02	UC2-ColonMetLiver
5408	562701	2066.N13.gz43 270599	F	M00055392D:A06	UC2-ColonMetLiver
5409	561457	2066,O13.gz43 270600	F	M00055394B;C06	UC2-ColonMetLiver
5410	549082	2066.C14.gz43 270604	F	M00055381A:F02	UC2-ColonMetLiver
5411	418622	2066.H14.gz43 270609	F	M00055385D:D03	UC2-ColonMetLiver
5412	238196	2066.K14.gz43 270612	F	M00055389A:D08	UC2-ColonMetLiver
5413	560868	2066.D15.gz43 270621	F	M00055381D:D08	UC2-ColonMetLiver
5414	459764	2066.E15.gz43 270622	F	M00055382D:D04	UC2-ColonMetLiver
5415	555639	2066.M15.gz43 270630	F	M00055391B:H08	UC2-ColonMetLiver
5416	556011	2066.O15.gz43 270632	F	M00055394B:D08	UC2-ColonMetLiver
5417	468565	2066.A17.gz43 270650	F	M00055379A:D07	UC2-ColonMetLiver
5418	448949	2066.B17.gz43 270651	F	M00055380A:C06	UC2-ColonMetLiver
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Table 2

	able 2				
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5419	636532	2066.J17.gz43 270659	F	M00055388B:E01	UC2-ColonMetLiver
5420	561487	2066.P17.gz43 270665	F	M00055395B;C04	UC2-ColonMetLiver
5421	50604	2066.L18.gz43 270677	F	M00055390C:E06	UC2-ColonMetLiver
5422	557249	2066,J20.gz43 270707	F	M00055388B:H04	UC2-ColonMetLiver
5423	562849	2066,H21.gz43 270721	F	M00055386A:G05	UC2-ColonMetLiver
5424	562386	2066.C22.gz43 270732	F	M00055381B:C10	UC2-ColonMetLiver
5425	635965	2066,J22.gz43_270739	F	M00055388C;D01	UC2-ColonMetLiver
5426	493487	2066.D23.gz43_270749	F	M00055382B;E02	UC2-ColonMetLiver
5427	44015	2066.N23.gz43_270759	F	M00055393B:F04	UC2-ColonMetLiver
5428	559854	2066.F24.gz43_270767	F	M00055383D:H11	UC2-ColonMctLiver
5429	282015	2066,K24,gz43_270772	F	M00055389C:F12	UC2-ColonMetLiver
5430	561206	2067.F02.gz43_270799	F	M00055402A;A05	UC2-ColonMetLiver
5431	486834	2067.G02.gz43_270800	F	M00055402D:A11	UC2-ColonMctLiver
5432	550730	2067.K02.gz43_270804	F	M00055407B:G07	UC2-ColonMetLiver
5433	560838	2067.P03.gz43_270825	F	M00055412D:A12	UC2-ColonMetLiver
5434	556382	2067.E04.gz43_270830	F	M00055400D:G01	UC2-ColonMetLiver
5435	561236	2067.I05.gz43_270850	F	M00055405A:A02	UC2-ColonMetLiver
5436	562584	2067.J05.gz43_270851	F	M00055406B:E07	UC2-ColonMetLiver
5437	550135	2067.N05.gz43_270855	F	M00055410C:G08	UC2-ColonMetLiver
5438	556380	2067.B06.gz43_270859	F	M00055397A;H07	UC2-ColonMetLiver
5439	511792	2067.C06.gz43_270860	F	M00055398C:F07	UC2-ColonMetLiver
5440	452761	2067.I06.gz43_270866	F	M00055405A:C01	UC2-ColonMctLiver
5441	465284	2067.K06.gz43_270868	F	M00055407C:E04	UC2-ColonMetLiver
5442	419489	2067,I07.gz43_270882	F	M00055405A:G11	UC2-ColonMetLiver
5443	540000	2067.P07.gz43_270889	F	M00055413A:A02	UC2-ColonMetLiver
5444	556105	2067.K08.gz43_270900	F	M00055407C:G04	UC2-ColonMetLiver
5445	562001	2067.N08.gz43_270903	F	M00055410C:H11	UC2-ColonMctLiver
5446	288626	2067.N09.gz43_270919	F	M00055410D:B02	UC2-ColonMetLiver
5447	561325	2067.H10.gz43_270929	F	M00055404A:B10	UC2-ColonMctLiver
5448	554742	2067.I10.gz43_270930	F	M00055405B:H05	UC2-ColonMctLiver
5449	560003	2067.A11.gz43_270938	F	M00055396B;C06	UC2-ColonMetLiver
5450	555856	2067.G11.gz43_270944	F	M00055403A;C07	UC2-ColonMetLiver
5451	633189	2067.N11.gz43_270951	F	M00055410D:G01	UC2-ColonMetLiver
5452	562147	2067.B12.gz43_270955	F	M00055397D;A01	UC2-ColonMetLiver
5453	553766	2067.H12.gz43_270961	F	M00055404A:D08	UC2-ColonMetLiver
5454	63669	2067.I13.gz43_270978	F	M00055405C:C04	UC2-ColonMetLiver
5455	491260	2067.J13.gz43_270979	F	M00055406D:A03	UC2-ColonMetLiver
5456	559776	2067.L13.gz43_270981	F	M00055408C:E04	UC2-ColonMetLiver
5457	561922	2067.A14.gz43_270986	F	M00055396B:G02	UC2-ColonMetLiver
5458	559102	2067.I14.gz43_270994	F	M00055405C:H07	UC2-ColonMetLiver
5459	440833	2067.014.gz43_271000	F	M00055412A:C05	UC2-ColonMetLiver
5460	561212	2067.K15.gz43_271012	F	M00055408A;A05	UC2-ColonMetLiver

Table 2

	able 2				
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ID			ORIE		
NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
5461	556734	2067.O15.gz43_271016	F	M00055412A:F09	UC2-ColonMetLiver
5462	487893	2067.J16.gz43_271027	F	M00055406D:C10	UC2-ColonMetLiver
5463	558180	2067.G18.gz43_271056	F	M00055403B:A10	UC2-ColonMetLiver
5464	557853	2067.J18.gz43 271059	F	M00055406D:G12	UC2-ColonMetLiver
5465	446171	2067.P19.gz43_271081	F	M00055413C:B09	UC2-ColonMetLiver
5466	561438	2067,B20.gz43_271083	F	M00055398A:C11	UC2-ColonMetLiver
5467	551617	2067.I20.gz43 271090	F	M00055405D:G05	UC2-ColonMetLiver
5468	469852	2067,B21.gz43 271099	F	M00055398B:A05	UC2-ColonMetLiver
5469	556126	2067.P21.gz43 271113	F	M00055413C:G09	UC2-ColonMetLiver
5470	551928	2067.F22.gz43 271119	F	M00055402C:C12	UC2-ColonMetLiver
5471	562236	2067.A23.gz43 271130	F	M00055397A:B10	UC2-ColonMetLiver
5472	456687	2067.B23.gz43 271131	F	M00055398B:C05	UC2-ColonMetLiver
5473	530715	2067.H23.gz43 271137	F	M00055404C:C11	UC2-ColonMetLiver
5474	66014	2067.K23.gz43_271140	F	M00055408A:F12	UC2-ColonMetLiver
5475	100821	2067.L23.gz43 271141	F	M00055409A:E06	UC2-ColonMetLiver
5476	154980	2067.A24.gz43 271146	F	M00055397A:C06	UC2-ColonMetLiver
5477	561236	2067.G24.gz43 271152	F	M00055403B:G12	UC2-ColonMetLiver
5478	516043	2067.L24.gz43 271157	F	M00055409A:E10	UC2-ColonMetLiver
5479	559380	2068.K01.gz43 271172	F	M00055424B:D04	UC2-ColonMetLiver
5480	449795	2068.O03.gz43 271208	F	M00055474A:G06	UC2-ColonMetLiver
5481	550166	2068.D04.gz43 271213	F	M00055417C:A05	UC2-ColonMetLiver
5482	561646	2068.F04.gz43 271215	F	M00055419D:D04	UC2-ColonMetLiver
5483	561112	2068.D05.gz43 271229	F	M00055417C:G03	UC2-ColonMetLiver
5484	418562	2068.E05.gz43 271230	F	M00055418C:D08	UC2-ColonMetLiver
5485	559385	2068.J05.gz43 271235	F	M00055423C;D05	UC2-ColonMetLiver
5486	561411	2068.M05.gz43 271238	F	M00055472A;B12	UC2-ColonMetLiver
5487	561770	2068.C07.gz43 271260	F	M00055416B:F11	UC2-ColonMetLiver
5488	557760	2068.D08.gz43 271277	F	M00055417D:D08	UC2-ColonMetLiver
5489	468672	2068.M08.gz43 271286	F	M00055472A:F02	UC2-ColonMetLiver
5490	556490	2068.N08.gz43 271287	F	M00055473B:D01	UC2-ColonMetLiver
5491	550018	2068.F09.gz43 271295	F	M00055419D:H07	UC2-ColonMetLiver
5492	561837	2068.H09.gz43 271297	F	M00055421C:F06	UC2-ColonMetLiver
5493	554028	2068.I09.gz43 271298	F	M00055422C:E10	UC2-ColonMetLiver
5494	470667	2068.D10.gz43 271309	F	M00055417D:H08	UC2-ColonMetLiver
5495	552561	2068.E10.gz43 271310	F	M00055418D:A03	UC2-ColonMetLiver
5496	556793	2068.G10,gz43 271312	F	M00055421A;A09	UC2-ColonMetLiver
5497	561718	2068.N10.gz43 271319	F	M00055473B:E10	UC2-ColonMetLiver
5498	630348	2068.E11.gz43 271326	F	M00055418D:B05	UC2-ColonMetLiver
5499	491728	2068.H11.gz43 271329	F	M00055421C:G07	UC2-ColonMetLiver
5500	562477	2068.C12.gz43 271340	F	M00055416D;B03	UC2-ColonMetLiver
5501	402836	2068.D12.gz43 271341	F	M00055418A:C12	UC2-ColonMetLiver
5502	556511	2068.J12.gz43 271347	F	M00055424A:A08	UC2-ColonMetLiver
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Table 2

	able 2				
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_	CLUSTER		NT	CLONE ID	LIBRARY
5503	559076	2068.N12.gz43_271351	F	M00055473C:A05	UC2-ColonMetLiver
5504	417426	2068.P12.gz43_271353	F	M00055475D:A05	UC2-ColonMetLiver
5505	456520	2068.A13.gz43_271354	F	M00055414C:C02	UC2-ColonMetLiver
5506	492982	2068.J13.gz43_271363	F	M00055424A;C10	UC2-ColonMetLiver
5507	558679	2068.C14.gz43_271372	F	M00055416D:D11	UC2-ColonMetLiver
5508	561707	2068.F14.gz43_271375	F	M00055420A:E12	UC2-ColonMetLiver
5509	561180	2068.I14.gz43_271378	F	M00055423A:A10	UC2-ColonMctLiver
5510	554953	2068.K14.gz43_271380	F	M00055425A:H03	UC2-ColonMetLiver
5511	492876	2068.L14.gz43_271381	F	M00055471C:B01	UC2-ColonMetLiver
5512	555933	2068.O14.gz43_271384	F	M00055474C:H12	UC2-ColonMetLiver
5513	562137	2068.P14.gz43_271385	F	M00055475D:A10	UC2-ColonMetLiver
5514	561500	2068.B15.gz43_271387	F	M00055415D:C12	UC2-ColonMetLiver
5515	561682	2068.D15.gz43_271389	F	M00055418A:E07	UC2-ColonMetLiver
5516	560118	2068.I15.gz43_271394	F	M00055423A:D02	UC2-ColonMetLiver
5517	512392	2068,J15.gz43_271395	F	M00055424A:F04	UC2-ColonMetLiver
5518	449356	2068.I16.gz43_271410	F	M00055423A:G08	UC2-ColonMetLiver
5519	447098	2068.L16.gz43 271413	F	M00055471C:D04	UC2-ColonMetLiver
5520	522220	2068.N16.gz43 271415	F	M00055473C:H05	UC2-ColonMetLiver
5521	453715	2068.P16.gz43_271417	F	M00055475D:B07	UC2-ColonMetLiver
5522	554774	2068.D17.gz43_271421	F	M00055418A:H04	UC2-ColonMetLiver
5523	553987	2068.I17.gz43_271426	F	M00055423B:A06	UC2-ColonMetLiver
5524	561645	2068.K17.gz43_271428	F	M00055425B:D12	UC2-ColonMetLiver
5525	549511	2068.M18.gz43_271446	F	M00055472C:G11	UC2-ColonMetLiver
5526	560700	2068.P18.gz43 271449	F	M00055475D:G08	UC2-ColonMetLiver
5527	493410	2068.K20.gz43 271476	F	M00055425B:F06	UC2-ColonMetLiver
5528	482425	2068.A21.gz43_271482	F	M00055414D:G10	UC2-ColonMetLiver
5529	561497	2068.B21.gz43 271483	F	M00055416A:C04	UC2-ColonMetLiver
5530	131130	2068.C21.gz43 271484	F	M00055417A:G11	UC2-ColonMetLiver
5531	524736	2068.F22.gz43 271503	F	M00055420C:E08	UC2-ColonMetLiver
5532	58680	2068.G22.gz43 271504	F	M00055421B:E05	UC2-ColonMetLiver
5533	562302	2068.N22.gz43 271511	F	M00055474A:B05	UC2-ColonMetLiver
5534	514824	2079.M01.gz43 271558	F	M00043021D:F10	UC3-NormColon
5535	452523	2079.P02.gz43 271577	F	M00043031A:B01	UC3-NormColon
5536	453572	2079.K04.gz43_271604	F	M00043012A:F06	UC3-NormColon
5537	406092	2079,I05,gz43 271618	F	M00043007D:F07	UC3-NormColon
5538	452759	2079.O05.gz43 271624	F	M00043029A:E11	UC3-NormColon
5539	456756	2079.B06.gz43 271627	F	M00042983D:A03	UC3-NormColon
5540	505226	2079.C06.gz43 271628	F	M00042987C:E04	UC3-NormColon
5541	238196	2079.D06.gz43 271629	F	M00042991C:C02	UC3-NormColon
5542	235874	2079.H06.gz43 271633	F	M00043005A:B06	UC3-NormColon
5543	512432	2079.O07.gz43 271656	F	M00043029B:C09	UC3-NormColon
5544	269927	2079.O09.gz43 271688	F	M00043029B:F09	UC3-NormColon

Table 2

	able 2				
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	CLUSTER	SEQ NAME		CLONE ID	LIBRARY
5545	449510	2079.K10.gz43_271700	F	M00043012D:A06	UC3-NormColon
5546	516799	2079.O10.gz43_271704	F	M00043029B:G10	UC3-NormColon
5547	455716	2079.E11.gz43_271710	F	M00042995C:E04	UC3-NormColon
5548	452976	2079.G11.gz43_271712	F	M00043003A:H07	UC3-NormColon
5549	452031	2079.N11.gz43_271719	F	M00043026D:D06	UC3-NormColon
5550	452957	2079.P11.gz43_271721	F	M00043032B:H01	UC3-NormColon
5551	452830	2079.D12.gz43_271725	F	M00042991D:F06	UC3-NormColon
5552	452220	2079.C13.gz43_271740	F	M00042988D:G10	UC3-NormColon
5553	500896	2079.K13.gz43_271748	F_	M00043013B:A06	UC3-NormColon
5554	455855	2079.L13.gz43_271749	F	M00043017D:B06	UC3-NormColon
5555	513381	2079.M13.gz43_271750	F	M00043023D:D06	UC3-NormColon
5556	448200	2079.B14.gz43_271755	F	M00042985B:C05	UC3-NormColon
5557	509027	2079.F14.gz43_271759	F	M00043000C:H06	UC3-NormColon
5558	452618	2079.N14.gz43_271767	F	M00043027B:C04	UC3-NormColon
5559	510254	2079.C15.gz43_271772	F	M00042989C:A06	UC3-NormColon
5560	451812	2079.D15.gz43_271773	F	M00042992B:B03	UC3-NormColon
5561	452392	2079.G15.gz43 271776	F	M00043003C:A02	UC3-NormColon
5562	504812	2079.I15.gz43 271778	F	M00043008D:E12	UC3-NormColon
5563	504431	2079.J15.gz43 271779	F	M00043011B:D03	UC3-NormColon
5564	452440	2079.L15.gz43 271781	F	M00043020A:A03	UC3-NormColon
5565	450225	2079.B16.gz43 271787	F	M00042985C:B03	UC3-NormColon
5566	511348	2079.C16.gz43 271788	F	M00042989C:B07	UC3-NormColon
5567	502343	2079.D16.gz43 271789	F	M00042992B:B11	UC3-NormColon
5568	451850	2079.F16.gz43 271791	F	M00043001A:C10	UC3-NormColon
5569	455117	2079.K16.gz43 271796	F	M00043013C:A01	UC3-NormColon
5570	452922	2079.L16.gz43 271797	F	M00043020A:G12	UC3-NormColon
5571	504501	2079.B17.gz43 27,1803	F	M00042985C:D02	UC3-NormColon
5572	503122	2079.F17.gz43_271807	F	M00043001B:C01	UC3-NormColon
5573	508126	2079.I17.gz43 271810	F	M00043008D:H09	UC3-NormColon
5574	448687	2079.A18.gz43 271818	F	M00042982C:G02	UC3-NormColon
5575	451491	2079.B18.gz43 271819	F	M00042985C:D04	UC3-NormColon
5576	452969	2079.F18.gz43 271823	F	M00043001B:E02	UC3-NormColon
5577	452866	2079.G18.gz43 271824	F	M00043003C:G10	UC3-NormColon
5578	446936	2079.K18.gz43 271828	F	M00043013D:F03	UC3-NormColon
5579	452445	2079.L18.gz43 271829	F	M00043021A:A09	UC3-NormColon
5580	451092	2079.N18.gz43 271831	F	M00043027D:G12	UC3-NormColon
5581	75212	2079.D19.gz43 271837	F	M00042992D:H02	UC3-NormColon
5582	452325	2079.F19.gz43 271839	F	M00043001B:H01	UC3-NormColon
5583	502984	2079,I19.gz43 271842	F	M00043009B;C09	UC3-NormColon
5584	508125	2079.J19.gz43 271843	F	M00043011C:H09	UC3-NormColon
5585	452528	2079.L19.gz43 271845	F	M00043021A:B06	UC3-NormColon
5586	509505	2079.C20.gz43 271852	F	M00042989D:A07	UC3-NormColon

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5589	453079	2079.M20.gz43_271862	F	M00043024B:H11	UC3-NormColon
5590	448813	2079.C21.gz43_271868	F	M00042990A:E05	UC3-NormColon
5591	513178	2079.D21.gz43_271869	F	M00042993A:D12	UC3-NormColon
5592	453804	2079.E21.gz43_271870	F	M00042996D:D12	UC3-NormColon
5593	453078	2079.L21.gz43_271877	F	M00043021A:H06	UC3-NormColon
5594	452611	2079.P21.gz43_271881	F	M00043033C:C05	UC3-NormColon
5595	504880	2079.A22.gz43_271882	F	M00042983A:E09	UC3-NormColon
5596	447445	2079.B22.gz43_271883	F	M00042986B:E06	UC3-NormColon
5597	503625	2079.G22.gz43_271888	F	M00043004A:C08	UC3-NormColon
5598	502614	2079.J22.gz43_271891	F	M00043011D:C12	UC3-NormColon
5599	454226	2079.K22.gz43_271892	F	M00043014B:G12	UC3-NormColon
5600	452973	2079.L22.gz43_271893	F	M00043021B:H03	UC3-NormColon
5601	456755	2079.M22.gz43_271894	F	M00043025A:C01	UC3-NormColon
5602	515127	2079.O22.gz43_271896	F	M00043030C:F03	UC3-NormColon
5603	507450	2079.F23.gz43_271903	F	M00043002A:G09	UC3-NormColon
5604	448381	2079.G23.gz43_271904	F	M00043004A:C10	UC3-NormColon
5605	505933	2079.K23.gz43_271908	F	M00043014C:F07	UC3-NormColon
5606	452648	2079.M23.gz43_271910	F	M00043025A:D09	UC3-NormColon
5607	453038	2079.N23.gz43 271911	F	M00043028C:H09	UC3-NormColon
5608	452611	2079.P23.gz43_271913	F	M00043033C:D05	UC3-NormColon
5609	501534	2079.B24.gz43_271915	F	M00042986C:B07	UC3-NormColon
5610	454849	2079.F24.gz43_271919	F	M00043002B:E06	UC3-NormColon
5611	366607	2079.H24.gz43_271921	F	M00043007B:D12	UC3-NormColon
5612	507066	2079.K24.gz43_271924	F	M00043014C:G06	UC3-NormColon
5613	519378	2090.B01.gz43_273908	F	M00043153B:B09	UC3-PrimColon
5614	453946	2090.H01.gz43_273914	F	M00043172C:G06	UC3-PrimColon
5615	527679	2090.L01.gz43_273918	F	M00043185B:D02	UC3-PrimColon
5616	454463	2090.P01.gz43_273922	F	M00043201D:D03	UC3-PrimColon
5617	521840	2090.D02.gz43_273926	F	M00043160B;E05	UC3-PrimColon
5618	528981	2090.M02.gz43_273935	F	M00043189C:F08	UC3-PrimColon
5619	528404	2090.C03.gz43_273941	F	M00043156C:E05	UC3-PrimColon
5620	523674	2090.N04.gz43_273968	F	M00043193C:G11	UC3-PrimColon
5621	446768	2090.F05.gz43_273976	F	M00043168C:F03	UC3-PrimColon
5622	128749	2090.H06.gz43_273994	F	M00043172D:H03	UC3-PrimColon
5623	525781	2090.K06.gz43_273997	F	M00043183C:B08	UC3-PrimColon
5624	452775	2090.N06.gz43_274000	F	M00043193C:H09	UC3-PrimColon
5625	524721	2090.L07.gz43_274014	F	M00043186C:A11	UC3-PrimColon
5626	454202	2090.M07.gz43_274015	F	M00043190A:B10	UC3-PrimColon
5627	452325	2090.J08.gz43_274028	F	M00043180C:B02	UC3-PrimColon
5628	453804	2090.F09.gz43_274040	F	M00043169A:F05	UC3-PrimColon

Table 2

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5629	454007	2090.O09.gz43_274049	F	M00043197D:H12	UC3-PrimColon
5630	425455	2090.L10.gz43_274062	F	M00043186D:B09	UC3-PrimColon
5631	453354	2090.H11.gz43_274074	F	M00043173B:F03	UC3-PrimColon
5632	525315	2090.I11.gz43_274075	F	M00043176C:A01	UC3-PrimColon
5633	453890	2090.N11.gz43_274080	F	M00043194A:G06	UC3-PrimColon
5634	454460	2090.A12.gz43_274083	F	M00043151B:D02	UC3-PrimColon
5635	522869	2090.C12.gz43_274085	F	M00043158A:F03	UC3-PrimColon
5636	454664	2090.I13.gz43_274107	F	M00043176C:E12	UC3-PrimColon
5637	528369	2090.L13.gz43_274110	F	M00043187C:E06	UC3-PrimColon
5638	526539	2090.J14.gz43_274124	F	M00043181C:C10	UC3-PrimColon
5639	452182	2090.O14.gz43_274129	F	M00043199A:F03	UC3-PrimColon
5640	520599	2090.H15.gz43_274138	F	M00043173C:D12	UC3-PrimColon
5641	530971	2090.L16.gz43_274158	F	M00043187D:H04	UC3-PrimColon
5642	289316	2090.N16.gz43_274160	F	M00043194D:E10	UC3-PrimColon
5643	454731	2090.A17.gz43_274163	F	M00043152A:F10	UC3-PrimColon
5644	453730	2090.D17.gz43_274166	F	M00043162D:E06	UC3-PrimColon
5645	450784	2090.G17.gz43_274169	F	M00043171D:G08	UC3-PrimColon
5646	404081	2090.M17.gz43_274175	F	M00043191C:G05	UC3-PrimColon
5647	456723	2090.A18.gz43_274179	F	M00043152A:G08	UC3-PrimColon
5648	526449	2090.C18.gz43_274181	F	M00043159A;C01	UC3-PrimColon
5649	413693	2090.H18.gz43_274186	F	M00043174B:B06	UC3-PrimColon
5650	454311	2090.J18.gz43_274188	F	M00043182B:C02	UC3-PrimColon
5651	526733	2090.H20.gz43_274218	F	M00043174C:C06	UC3-PrimColon
5652	432159	2090.K20.gz43_274221	F	M00043184D:G06	UC3-PrimColon
5653	453983	2090.M20.gz43_274223	F	M00043192B;H06	UC3-PrimColon
5654	518172	2090.N20.gz43_274224	F	M00043195D:A05	UC3-PrimColon
5655	454629	2090.P22.gz43_274258	F	M00043210C:E08	UC3-PrimColon
5656	530939	2090.A23.gz43_274259	F	M00043152B:H04	UC3-PrimColon
5657	234270	2090.G23.gz43_274265	F	M00043172C:B05	UC3-PrimColon
5658	529742	2090.I24.gz43_274283	F	M00043178C:G06	UC3-PrimColon
5659	404461	2092.G04.gz43_274444	F	M00043438B:E10	UC3-ColonMetLiver
5660	451023	2092.I04.gz43_274446	F	M00043448D:H09	UC3-ColonMetLiver
5661	535436	2092.G05.gz43_274460	F	M00043438C:D12	UC3-ColonMetLiver
5662	456001	2092.N05.gz43_274467	F	M00043470C:A01	UC3-ColonMetLiver
5663	450724	2092.O06.gz43_274484	F	M00043478B:D01	UC3-ColonMetLiver
5664	539142	2092.A07.gz43_274486	F	M00042592A:H02	UC3-ColonMetLiver
5665	450914	2092.K07.gz43_274496	F	M00043455C:G07	UC3-ColonMetLiver
5666	455855	2092.P07.gz43_274501	F	M00043484D:H08	UC3-ColonMetLiver
5667	537451	2092.F08.gz43_274507	F	M00043433A:F02	UC3-ColonMetLiver
5668	535123	2092.G08.gz43_274508	F	M00043439C:D05	UC3-ColonMetLiver
5669	452830	2092.J08.gz43_274511	F	M00043452D:D05	UC3-ColonMetLiver
5670	534054	2092.L08.gz43_274513	F	M00043460D:C03	UC3-ColonMetLiver

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Table 2

	able 2				
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5671	162851	2092.M08.gz43_274514	F	M00043465B:G08	UC3-ColonMetLiver
5672	455289	2092.D09.gz43_274521	F	M00043420C:C02	UC3-ColonMetLiver
5673	456020	2092.H09.gz43_274525	F	M00043446B:H01	UC3-ColonMetLiver
5674	448946	2092.I09.gz43_274526	F	M00043449C:H09	UC3-ColonMetLiver
5675	539353	2092.L09.gz43_274529	F	M00043460D:H01	UC3-ColonMetLiver
5676	450658	2092.P09.gz43_274533	F	M00043485A;C04	UC3-ColonMetLiver
5677	544355	2092.E10.gz43_274538	F	M00043427C:E11	UC3-ColonMetLiver
5678	452128	2092.F10.gz43_274539	F	M00043434A:H02	UC3-ColonMetLiver
5679	447534	2092.I10.gz43_274542	F	M00043449D:A06	UC3-ColonMetLiver
5680	455256	2092.P10.gz43_274549	F	M00043485B:C09	UC3-ColonMetLiver
5681	455201	2092.C11.gz43 274552	F	M00042640C:C10	UC3-ColonMetLiver
5682	535129	2092.F11.gz43 274555	F	M00043434B;D06	UC3-ColonMetLiver
5683	455808	2092.H11.gz43_274557	F	M00043446C:E04	UC3-ColonMetLiver
5684	215366	2092.I11.gz43 274558	F	M00043449D:E09	UC3-ColonMetLiver
5685	456254	2092.J11.gz43 274559	F	M00043453B:C06	UC3-ColonMetLiver
5686	536415	2092.N11.gz43_274563	F	M00043472A:E06	UC3-ColonMetLiver
5687	452801	2092.F12.gz43 274571	F	M00043434B:E04	UC3-ColonMetLiver
5688	400258	2092.H12.gz43 274573	F	M00043446C:E08	UC3-ColonMetLiver
5689	546121	2092.J12.gz43 274575	F	M00043453B:F02	UC3-ColonMetLiver
5690	454621	2092.K12.gz43 274576	F	M00043457C:B12	UC3-ColonMetLiver
5691	455162	2092.D13.gz43 274585	F	M00043422B:C03	UC3-ColonMetLiver
5692	543429	2092.P13.gz43 274597	F	M00043486B:D02	UC3-ColonMetLiver
5693	456249	2092,I14,gz43 274606	F	M00043450B:C11	UC3-ColonMetLiver
5694	531461	2092.E15.gz43_274618	F	M00043428C:A08	UC3-ColonMetLiver
5695	455572	2092.H15.gz43 274621	F	M00043447A;F02	UC3-ColonMetLiver
5696	456783	2092.I15.gz43 274622	F	M00043450B:H05	UC3-ColonMetLiver
5697	2544	2092.N15.gz43 274627	F	M00043473C:B11	UC3-ColonMetLiver
5698	543431	2092.P15.gz43 274629	F	M00043486C:D02	UC3-ColonMetLiver
5699	402534	2092.O16.gz43 274644	F	M00043481A:G02	UC3-ColonMetLiver
5700	454563	2092.I17.gz43 274654	F	M00043450C;F11	UC3-ColonMetLiver
5701	541901	2092.K17.gz43_274656	F	M00043459A:B08	UC3-ColonMetLiver
5702	536225	2092.L17.gz43 274657	F	M00043462C:E12	UC3-ColonMetLiver
5703	455379	2092.M17.gz43 274658	F	M00043468C:D08	UC3-ColonMetLiver
5704	456567	2092.P17.gz43 274661	F	M00043486C:F07	UC3-ColonMetLiver
5705	539804	2092.C18.gz43 274664	F	M00043417A:H12	UC3-ColonMetLiver
5706	452618	2092.C18.gz43_274604 2092.N18.gz43_274675	F	M00043477A;H12	UC3-ColonMetLiver
5707	543855	2092.J19.gz43 274687	F	M00043473D:D11	UC3-ColonMetLiver
5708	415326	2092.L19.gz43 274689	F	M00043453D;D02	UC3-ColonMetLiver
5709	455117	2092.B20.gz43_274689	F	M00043402D:C09	UC3-ColonMetLiver
5710	455601	2092.C20.gz43 274696	F	M00042000D:B08	UC3-ColonMetLiver
5711	455254	2092.E20.gz43_274698	F	M00043417B:F12	UC3-ColonMetLiver
5712	402916	2092.E20.gz43_274698 2092.I20.gz43_274702	F	M00043451A:B02	UC3-ColonMetLiver
3/12	402910	2072.120.gz43_2/4/02	г	1V100043431A;B02	ocs-colonivietraver

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5713	455864	2092.N20.gz43_274707	F	M00043474C:H05	UC3-ColonMetLiver
5714	27586	2092.P20.gz43_274709	F	M00043489C:H06	UC3-ColonMetLiver
5715	452392	2092.B21.gz43_274711	F	M00042600D:E11	UC3-ColonMetLiver
5716	447380	2092.C21.gz43_274712	F	M00043417C:F12	UC3-ColonMetLiver
5717	448250	2092.I21.gz43_274718	F	M00043451B:D08	UC3-ColonMetLiver
5718	536171	2092.B22.gz43_274727	F	M00042637C:E12	UC3-ColonMetLiver
5719	455501	2092.H22.gz43_274733	F	M00043448B:E08	UC3-ColonMetLiver
5720	455808	2092.O22.gz43_274740	F	M00043483B:G01	UC3-ColonMctLiver
5721	422223	2092.C23.gz43_274744	F	M00043418A:A06	UC3-ColonMetLiver
5722	455814	2092.023.gz43_274756	F	M00043483B:G11	UC3-ColonMetLiver
5723	538582	2092.H24.gz43_274765	F	M00043448C:G07	UC3-ColonMetLiver
5724	447238	2092.I24.gz43_274766	F	M00043451C:H03	UC3-ColonMetLiver
5725	456577	2092.P24.gz43 274773	F	M00043490A:F03	UC3-ColonMetLiver
5726	11576	2176.F01.gz43 281685	F	M00039503B:G05	HMEC-Unt
5727	401426	2176.P01.gz43 281695	F	M00039605D:E02	- HMEC-Unt
5728	409330	2176.B02.gz43 281697	F	M00039363B:H10	HMEC-Unt
5729	407195	2176.C02.gz43 281698	F	M00039369D:D11	HMEC-Unt
5730	411985	2176.D02.gz43 281699	F	M00039379A:D03	HMEC-Unt
5731	412621	2176.F02.gz43 281701	F	M00039503D:E08	HMEC-Unt
5732	407711	2176.H02.gz43 281703	F	M00039514B:E07	HMEC-Unt
5733	411113	2176.G03.gz43 281718	F	M00039508D:C03	HMEC-Unt
5734	402411	2176.I04.gz43 281736	F	M00039523A:D09	HMEC-Unt
5735	147279	2176.F06.gz43 281765	F	M00039504B:F05	HMEC-Unt
5736	400628	2176.L06.gz43 281771	F	M00039566D:E08	HMEC-Unt
5737	407443	2176.C07.gz43 281778	F	M00039371A:E01	HMEC-Unt
5738	413346	2176.N07.gz43 281789	F	M00039592C:F09	HMEC-Unt
5739	125543	2176.O07.gz43 281790	F	M00039599A:E08	HMEC-Unt
5740	202308	2176.P07.gz43 281791	F	M00039606C:B07	HMEC-Unt
5741	400428	2176.K08.gz43 281802	F	M00039562A:D10	HMEC-Unt
5742	400426	2176.H09.gz43 281815	F	M00039516C:H06	HMEC-Unt
5743	402147	2176.P09.gz43 281823	F	M00039607A:F05	HMEC-Unt
5744	400233	2176.K10.gz43 281834	F	M00039562D:B02	HMEC-Unt
5745	408986	2176.K11.gz43 281850	F	M00039562D:G01	HMEC-Unt
5746	400171	2176.B12.gz43 281857	F	M00039366C:B10	HMEC-Unt
5747	403671	2176.C12.gz43 281858	F	M00039373B:B09	HMEC-Unt
5748	400047	2176.A13.gz43 281872	F	M00039184A:A08	HMEC-Unt
5749	404736	2176.B13.gz43 281873	F	M00039366D:A02	HMEC-Unt
5750	413627	2176.F13.gz43 281877	F	M00039505B:G07	HMEC-Unt
5751	407260	2176.K13.gz43 281882	F	M00039563C:D01	HMEC-Unt
5752	155374	2176.M13.gz43 281884	F	M00039586A:C03	HMEC-Unt
5753	409612	2176.I14.gz43 281896	F	M00039525B:H03	HMEC-Unt
5754	402411	2176.A16,gz43 281920	F	M00039184C:C05	HMEC-Unt
2,54	.02711	21. 0.1110.g2+3_201920	لـــُـا	1.1000371070.003	THILL OIL

Table 2

	able 2				
SEQ					
ID			ORIE		
ИО	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
5755	411124	2176.C17.gz43_281938	F	M00039375A:C09	HMEC-Unt
5756	412416	2176.J17.gz43_281945	F	M00039534D:E07	HMEC-Unt
5757	406734	2176.M17.gz43_281948	F	M00039586D:D05	HMEC-Unt
5758	408306	2176.K18.gz43_281962	F	M00039564B:F08	HMEC-Unt
5759	1054	2176.E19.gz43_281972	F	M00039502D:H03	HMEC-Unt
5760	406855	2176.I19.gz43_281976	F	M00039527B:D06	HMEC-Unt
5761	400454	2176.H20.gz43_281991	F	M00039521A:D02	HMEC-Unt
5762	402147	2176.J20.gz43_281993	F	M00039536B:E10	HMEC-Unt
5763	202308	2176.M20.gz43_281996	F	M00039588A:H12	HMEC-Unt
5764	413997	2176.J21.gz43_282009	F	M00039536B:H03	HMEC-Unt
5765	409589	2176.K21.gz43_282010	F	M00039564C:H05	HMEC-Unt
5766	409262	2176.M21.gz43_282012	F	M00039588B:H02	HMEC-Unt
5767	405932	2176.H22.gz43_282023	F	M00039521C:B01	HMEC-Unt
5768	20517	2176.M22.gz43_282028	F	M00039588D:F10	HMEC-Unt
5769	413915	2176.P23.gz43_282047	F	M00039610A:G11	HMEC-Unt
5770	412483	2176.F24.gz43_282053	F	M00039507D:E03	HMEC-Unt
5771	402392	2176.G24.gz43_282054	F	M00039514A:G03	HMEC-Unt
5772	401510	2176.J24.gz43_282057	F	M00039537B;F06	HMEC-Unt
5773	408649	2176.L24.gz43_282059	F	M00039583A:G09	HMEC-Unt
5774	418682	2178.K02.gz43_282160	F	M00040183B:C06	HMEC-bFGF
5775	417259	2178.E04.gz43_282186	F	M00040141A:G10	HMEC-bFGF
5776	20517	2178.P04.gz43_282197	F	M00040235C:D02	HMEC-bFGF
5777	414821	2178.G05.gz43_282204	F	M00040160C:A04	HMEC-bFGF
5778	402534	2178.K05.gz43_282208	F	M00040183D:C08	HMEC-bFGF
5779	402411	2178.D07.gz43_282233	F	M00040134A:A07	HMEC-bFGF
5780	419255	2178.L07.gz43_282241	F	M00040188B:E05	HMEC-bFGF
5781	417426	2178.G11.gz43_282300	F	M00040161C:H06	HMEC-bFGF
5782	415527	2178.G12.gz43_282316	F	M00040161D:C03	HMEC-bFGF
5783	418340	2178.J12.gz43_282319	F	M00040181B:B06	HMEC-bFGF
5784	403154	2178.L12.gz43_282321	F	M00040189D;A06	HMEC-bFGF
5785	403837	2178.J13.gz43_282335	F	M00040181B:C05	HMEC-bFGF
5786	418482	2178.L14.gz43_282353	F	M00040190B:C02	HMEC-bFGF
5787	419751	2178.C15.gz43_282360	F	M00039760B:F12	HMEC-bFGF
5788	402070	2178.B16.gz43_282375	F	M00039752D:D07	HMEC-bFGF
5789	402070	2178.N16.gz43_282387	F	M00040227A:E07	HMEC-bFGF
5790	420504	2178.C18.gz43_282408	F	M00039760C:H07	HMEC-bFGF
5791	147279	2178.P19.gz43_282437	F	M00040248A:G09	HMEC-bFGF
5792	402353	2178.H20.gz43_282445	F	M00040171B:H03	HMEC-bFGF
5793	416914	2178.L20.gz43_282449	F	M00040196B:F10	HMEC-bFGF
5794	163970	2178.E21.gz43_282458	F	M00040145B:C12	HMEC-bFGF
5795	416762	2178.O21.gz43_282468	F	M00040233C:F09	HMEC-bFGF
5796	57183	2178.M22.gz43 282482	F	M00040222A:E06	HMEC-bFGF

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T	able 2				
SEQ					
ID			ORIE		
NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
5797	403306	2184.D01.gz43_282670	F	M00039950C:C05	HMEC-VEGF
5798	402298	2184.L01.gz43_282678	F	M00040328D:A03	HMEC-VEGF
5799	420958	2184.K02.gz43_282693	F	M00040322B:A08	HMEC-VEGF
5800	423884	2184.A03.gz43 282699	F	M00039747A:H06	HMEC-VEGF
5801	268336	2184.J04.gz43 282724	F	M00040318C:B09	HMEC-VEGF
5802	422590	2184.O07.gz43_282777	F	M00040364B:E11	HMEC-VEGF
5803	402298	2184.A08.gz43 282779	F	M00039749B:G05	HMEC-VEGF
5804	423534	2184.J11.gz43 282836	F	M00040319D:G10	HMEC-VEGF
5805	423008	2184.K13.gz43 282869	F	M00040323D:F04	HMEC-VEGF
5806	424723	2184.N13.gz43 282872	F	M00040347B:B11	HMEC-VEGF
5807	20517	2184.D16.gz43 282910	F	M00039958D:D05	HMEC-VEGF
5808	403306	2184.A17.gz43 282923	F	M00039750C:F08	HMEC-VEGF
5809	140224	2184.A22.gz43 283003	F	M00039762C:D11	HMEC-VEGF
5810	402298	2184.G24.gz43_283041	F	M00040294B:E09	HMEC-VEGF
5811	468930	1564.A01.gz43_296552	F	M00042709B:G05	UC2-NormColon
5812	463143	1564.B01.gz43 296553	F	M00042711B:H04	UC2-NormColon
5813		1564.C01.gz43 296554	F	M00042713B:C08	UC2-NormColon
5814		1564.I01.gz43_296560	F	M00042724A:G02	UC2-NormColon
5815		1564.J01.gz43_296561	F	M00042726D:G11	UC2-NormColon
5816		1564.O01.gz43 296566	F	M00042737B:C07	UC2-NormColon
5817		1564.C03.gz43 296586	F	M00042737B:E07	UC2-NormColon
5818		1564.H03.gz43 296591	F	M00042713B:F03	UC2-NormColon
5819	446740	1564.004.gz43_296614	F	M00042722A:G08	UC2-NormColon
5820	447268	1564.C05.gz43 296618	F	M00042713C:B08	UC2-NormColon
5821		1564.K05.gz43 296626	F	M00042713C:B08	UC2-NormColon
5822	420686	1564.N05.gz43 296629	F	M00042725A:F11	UC2-NormColon
5823	447579	1564.O05.gz43_296630	F	M00042737C:E03	UC2-NormColon
5824	447006	1564.B06.gz43 296633	F	M00042737C.E03	UC2-NormColon
5825	463824	1564.N06.gz43 296645	F	M00042711C:H10	UC2-NormColon UC2-NormColon
5826		1564.G07.gz43 296654	F	M00042733D:A07	UC2-NormColon
5827		1564.A08.gz43 296664	F	M00042720C:E03	UC2-NormColon
5828			F		
5829	462986 454825	1564.K08.gz43_296674 1564.M08.gz43_296676	F	M00042729A:H08	UC2-NormColon
			F	M00042733C:C05	UC2-NormColon
5830	466614	1564.O08.gz43 296678	F	M00042737D:D03	UC2-NormColon
5831	457975	1564.A09.gz43_296680	F	M00042710B:B09	UC2-NormColon
5832	446345	1564.I09.gz43_296688		M00042724D:B04	UC2-NormColon
5833		1564.C11.gz43_296714	F	M00042714A:G04	UC2-NormColon
5834		1564.E11.gz43 296716	F	M00042716D:G04	UC2-NormColon
5835	447387	1564.N11.gz43_296725	F	M00042736A:F03	UC2-NormColon
5836		1564.B12.gz43_296729	F	M00042712A:E08	UC2-NormColon
5837	447561	1564.C12.gz43_296730	F	M00042714A:H05	UC2-NormColon
5838	461313	1564.K12.gz43 296738	F	M00042729B:F10	UC2-NormColon

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Table 2

1	able 2				
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ID			ORIE		
NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
5839	453679	1564.L12.gz43 296739	F	M00042732A:A11	UC2-NormColon
5840	446964	1564.G13.gz43_296750	F	M00042721A:G07	UC2-NormColon
5841	459158	1564.H13.gz43_296751	F	M00042722D:C12	UC2-NormColon
5842	450723	1564.M13.gz43_296756	F	M00042733D:G08	UC2-NormColon
5843	447826	1564.C15.gz43_296778	F	M00042714B:E01	UC2-NormColon
5844	446741	1564.D15.gz43_296779	F	M00042715D:E02	UC2-NormColon
5845	452687	1564.F15.gz43_296781	F	M00042719A:H06	UC2-NormColon
5846	446601	1564.H15.gz43_296783	F	M00042723A:D09	UC2-NormColon
5847	447649	1564.B19.gz43_296841	F	M00042712C:E06	UC2-NormColon
5848	465528	1564.C19.gz43_296842	F	M00042714C:C10	UC2-NormColon
5849	462865	1564.E19.gz43 296844	F	M00042717B:G11	UC2-NormColon
5850	405932	1564.F19.gz43_296845	F	M00042719C:H05	UC2-NormColon
5851	457922	1564.G19.gz43_296846	F	M00042721D:B03	UC2-NormColon
5852	463821	1564.N19.gz43 296853	F	M00042737A:A07	UC2-NormColon
5853	463217	1564.O19.gz43 296854	F	M00042738D:H12	UC2-NormColon
5854	389425	1564.A20.gz43 296856	F	M00042711A:F01	UC2-NormColon
5855	446242	1564.O20.gz43 296870	F	M00042739A:C06	UC2-NormColon
5856	449171	1564.C21.gz43 296874	F	M00042714C:F12	UC2-NormColon
5857	450723	1564.G21.gz43 296878	F	M00042721D:D01	UC2-NormColon
5858	469766	1564.M21.gz43 296884	F	M00042735A:G12	UC2-NormColon
5859	446866	1564.D22.gz43 296891	F	M00042716A:F09	UC2-NormColon
5860	403949	1564.M23.gz43_296916	F	M00042735B:A06	UC2-NormColon
5861	446922	1564.O23.gz43_296918	F	M00042739A:G07	UC2-NormColon
5862	235874	1564.C24.gz43_296922	F	M00042714D:E11	UC2-NormColon
5863	558301	1681.A01.gz43_296936	F	M00055739A:C09	UC2-NormColon
5864	426366	1681.I01.gz43_296944	F	M00055749D:B07	UC2-NormColon
5865	638943	1681.J01.gz43_296945	F	M00055750C:H10	UC2-NormColon
5866	645603	1681.O01.gz43_296950	F	M00055758C:H10	UC2-NormColon
5867	642644	1681.E02.gz43_296956	F	M00055744C:D02	UC2-NormColon
5868	644105	1681.I02.gz43_296960	F	M00055749D:C01	UC2-NormColon
5869	468613	1681.A03.gz43_296968	F	M00055739B:B06	UC2-NormColon
5870	648905	1681.B03.gz43_296969	F	M00055740C;A07	UC2-NormColon
5871	446925	1681.H03.gz43_296975	F	M00055748C:C07	UC2-NormColon
5872	649356	1681.I03.gz43_296976	F	M00055749D:D06	UC2-NormColon
5873	510724	1681.K03.gz43_296978	F	M00055752C;C06	UC2-NormColon
5874	446676	1681.L03.gz43_296979	F	M00055753D:C06	UC2-NormColon
5875	644937	1681.B04.gz43_296985	F	M00055740C:E06	UC2-NormColon
5876	644342	1681.I04.gz43_296992	F	M00055749D:F12	UC2-NormColon
5877	640264	1681.004.gz43 296998	F	M00055758D:E02	UC2-NormColon
5878	463951	1681.D05.gz43_297003	F	M00055743C:D12	UC2-NormColon
5879	644345	1681.K05.gz43_297010	F	M00055752C:F06	UC2-NormColon
5880	646552	1681.N06.gz43 297029	F	M00055757B:B07	UC2-NormColon

Table 2

	able 2				
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	CLUSTER		NT	CLONE ID	LIBRARY
5881	611927	1681.K07.gz43_297042	F	M00055752C:H07	UC2-NormColon
5882	639420	1681.N07.gz43_297045	F	M00055757B:C04	UC2-NormColon
5883	568467	1681.A08.gz43_297048	F	M00055739B;H08	UC2-NormColon
5884	645147	1681.C08.gz43_297050	F	M00055742B:H06	UC2-NormColon
5885	447936	1681.E08.gz43_297052	F	M00055744D:A11	UC2-NormColon
5886	492917	1681.O08.gz43_297062	F	M00055759A:B02	UC2-NormColon
5887	649427	1681.D09.gz43_297067	F	M00055743D:D04	UC2-NormColon
5888	556654	1681.F09.gz43_297069	F	M00055745D:F11	UC2-NormColon
5889	594040	1681.H09.gz43_297071	F	M00055748D:C03	UC2-NormColon
5890	488432	1681.J09.gz43_297073	F	M00055751A:F06	UC2-NormColon
5891	641144	1681.K09.gz43_297074	F	M00055752D:C01	UC2-NormColon
5892	550223	1681.L09.gz43_297075	F	M00055754A:E07	UC2-NormColon
5893	491544	1681.M09.gz43_297076	F	M00055755D:C09	UC2-NormColon
5894	648481	1681.A10.gz43_297080	F	M00055739C:D11	UC2-NormColon
5895	150839	1681.B10.gz43_297081	F	M00055740D:G12	UC2-NormColon
5896	644053	1681.F10.gz43_297085	F	M00055746A:C09	UC2-NormColon
5897	644342	1681.K10.gz43_297090	F	M00055752D:F01	UC2-NormColon
5898	419479	1681.C11.gz43_297098	F	M00055742C:C01	UC2-NormColon
5899	645000	1681.E11.gz43_297100	F	M00055744D:F03	UC2-NormColon
5900	478833	1681.L11.gz43_297107	F	M00055754A:H06	UC2-NormColon
5901	644684	1681.A13.gz43_297128	F	M00055739D:B12	UC2-NormColon
5902	640504	1681.B13.gz43_297129	F	M00055741A:D09	UC2-NormColon
5903	463815	1681.A14.gz43_297144	F	M00055739D:C03	UC2-NormColon
5904	640997	1681.B15.gz43_297161	F	M00055741B:B12	UC2-NormColon
5905	642263	1681.D15.gz43_297163	F	M00055744A:B04	UC2-NormColon
5906	451508	1681.P15.gz43_297175	F	M00055761A:E06	UC2-NormColon
5907	644919	1681.A16.gz43_297176	F	M00055739D:E04	UC2-NormColon
5908	639703	1681.K16.gz43_297186	F	M00055753A:D04	UC2-NormColon
5909	639750	1681.M16.gz43_297188	F	M00055756C:C03	UC2-NormColon
5910	587696	1681.F17.gz43_297197	F	M00055746C:F06	UC2-NormColon
5911	461917	1681.H17.gz43_297199	F	M00055749B:C10	UC2-NormColon
5912	644510	1681,I17,gz43_297200	F	M00055750B:H01	UC2-NormColon
5913	526459	1681.B18.gz43_297209	F	M00055741C:A09	UC2-NormColon
5914	471277	1681.F18.gz43_297213	F	M00055746C:F10	UC2-NormColon
5915	483042	1681.K18.gz43_297218	F	M00055753B:A02	UC2-NormColon
5916	514838	1681.L18.gz43_297219	F	M00055755A:B11	UC2-NormColon
5917	643984	1681.N18.gz43_297221	F	M00055757D:B05	UC2-NormColon
5918	462245	1681.A19.gz43_297224	F	M00055740A:B03	UC2-NormColon
5919	640017	1681.D19.gz43_297227	F	M00055744B:B02	UC2-NormColon
5920	453726	1681.E19.gz43_297228	F	M00055745A:H02	UC2-NormColon
5921	644468	1681.F19.gz43_297229	F	M00055746C:G06	UC2-NormColon
5922	644105	1681.H19.gz43 297231	F	M00055749C:B03	UC2-NormColon
			لـــتـــا		

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Table 2

able 2				
		one		
CY Y CORPA	CTC 3713 CT			
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				UC2-NormColon
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		_		UC2-NormColon
				UC2-NormColon
				UC2-NormColon
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			M00055745B:H02	UC2-NormColon
	1681.J23.gz43_297297		M00055752A:G10	UC2-NormColon
463824	1681.A24.gz43_297304		M00055740B:F09	UC2-NormColon
389377	1681.C24.gz43_297306	F	M00055743B:E01	UC2-NormColon
468959	1681.E24.gz43_297308	F	M00055745C:A06	UC2-NormColon
553675	1681.M24.gz43_297316	F	M00055757A:A07	UC2-NormColon
449978	1561.C01.gz43_314395	F	M00042344D:F03	UC2-NormColon
463920	1561.G01.gz43_314399	F	M00042526D:A05	UC2-NormColon
456267	1561.J01.gz43_314402	F	M00042531B:G12	UC2-NormColon
459997	1561.M01.gz43_314405	F	M00042537A:D12	UC2-NormColon
447386	1561.I02.gz43_314417	F	M00042529D:D07	UC2-NormColon
468257	1561.K02.gz43_314419	F	M00042533B:F11	UC2-NormColon
18786	1561.E03.gz43 314429	F	M00042523C:H06	UC2-NormColon
447494	1561.A04.gz43_314441	F	M00042341A:D08	UC2-NormColon
459893	1561.B04.gz43_314442	F	M00042343B:D05	UC2-NormColon
462558	1561.M04.gz43_314453	F	M00042537A:G09	UC2-NormColon
458425	1561.O04.gz43_314455	F	M00042541A;B07	UC2-NormColon
249178	1561.C05.gz43 314459	F	M00042345B;A05	UC2-NormColon
462815	1561.J05.gz43 314466	F	M00042531D:G08	UC2-NormColon
417291	1561.K05.gz43 314467	F	M00042533C:F04	UC2-NormColon
463341	1561.M05.gz43 314469	F	M00042537A:H04	UC2-NormColon
447868	1561.A06.gz43 314473	F	M00042341A:H04	UC2-NormColon
461653		F		UC2-NormColon
465127		F	M00042533D:B05	UC2-NormColon
460246	1561.N06.gz43 314486	F	M00042539B:D09	UC2-NormColon
33139		F	M00042541A:E06	UC2-NormColon
457508		F		UC2-NormColon
461917		_		UC2-NormColon
				UC2-NormColon
				UC2-NormColon
		-		UC2-NormColon
				UC2-NormColon
		-		UC2-NormColon
		$\overline{}$		UC2-NormColon
	CLUSTER 447485 218416 489040 649702 643142 448251 644461 641968 470667 644479 463824 338377 468959 553675 549978 447386 447494 458257 18786 447494 459893 462558 458425 249178 462815 417291 463341 447868 46165127 460246 33139	CLUSTER SEQ NAME 447485 1681.K19.gz43 297234 218416 1681.D20.gz43 297243 489040 1681.H20.gz43 297249 649702 1681.H20.gz43 297249 643142 1681.A21.gz43 297269 643143 1681.A21.gz43 297269 644161 1681.N21.gz43 297269 644168 1681.R21.gz43 297269 644168 1681.R21.gz43 297269 644168 1681.R21.gz43 297269 644169 1681.R22.gz43 297209 644479 1681.L23.gz43 297304 648959 1681.E23.gz43 297306 648959 1681.E24.gz43 297316 648978 1561.C01.gz43 314497 64978 1561.M01.gz43 314402 645267 1561.J01.gz43 314402 645267 1561.J01.gz43 314402 64527 1561.K02.gz43 314419 18786 1561.E03.gz43 314419 18786 1561.E03.gz43 314442 642558 1561.M04.gz43 314442 642558 1561.M04.gz43 314442 642558 1561.M04.gz43 314443 646245 1561.S05.gz43 314466 647729 1 561.K02.gz43 314466 65127 1561.K02.gz43 314466	CLUSTER SEQ NAME NT 447485 1681.K19.gz43 297234 F 489040 1681.D20.gz43 297234 F 649702 1681.D20.gz43 297234 F 649702 1681.D20.gz43 297234 F 649702 1681.D20.gz43 297236 F 643142 1681.A21.gz43 297256 F 644461 1681.N21.gz43 297260 F 644461 1681.N21.gz43 297267 F 644461 1681.N21.gz43 297267 F 644461 1681.N21.gz43 297267 F 644491 1681.D21.gz43 297292 F 644479 1681.D23.gz43 297292 F 644479 1681.D23.gz43 297304 F 839377 1681.C24.gz43 297306 F 648959 1681.E24.gz43 297306 F 648959 1681.E24.gz43 297306 F 648997 1561.D01.gz43 314395 F 649978 1561.C01.gz43 314402 F 649978 1561.K02.gz43 314417 F 646257 1561.M01.gz43 314417 F 646257 1561.M02.gz43 314419 F 18786 1561.E02.gz43 314419 F 18786 1561.E02.gz43 314417 F 646258 1561.M04.gz43 314442 F 645258 1561.M04.gz43 314442 F 646258 1561.M04.gz43 314442 F 646258 1561.M04.gz43 314447 F 646258 1561.M04.gz43 314447 F 646251 1561.O5.gz43 314466 F 647794 1561.K05.gz43 314467 F 6463341 1561.K05.gz43 314467 F 646341 1561.K05.gz43 314468 F 647704 1561.K05.gz43 314467 F 647704 1561.K05.gz43 314468 F 6477054 1561.K05.gz43 314467 F 6477064 1561.K05.gz43 314468 F 6477064 1561.K05.gz43 314468 F 6477064 1561.K05.gz43 314468 F 6477064 1561.K05.gz43 314468 F 6477064 1561.K05.gz43 314487 F 6477064 1561.K05.gz43	CLUSTER SEQ NAME

Table 2

DRI		able 2				
NO CLUSTER SEQ NAME NT CLONE ID LIBRARY	SEQ					
Section						
S966	NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
5967 464510 1561.D11.g243 314556 F M0004252D-A08 UC2-NormColon 5968 457846 1561.G11.g243 314559 F M0004252D-A08 UC2-NormColon 5969 15296 1561.H12.g243 314571 F M00042330C-H02 UC2-NormColon 5970 446578 1561.H12.g243 314571 F M0004235D-D04 UC2-NormColon 5971 464287 1561.H12.g243 314577 F M0004253D-D09 UC2-NormColon 5973 446183 1561.G14.g243 314601 F M0004253C-C101 UC2-NormColon 5974 446183 1561.G14.g243 314609 F M0004253C-C101 UC2-NormColon 5975 468565 1561.H4.g243 314609 F M0004253C-C10 UC2-NormColon 5976 470600 1561.J14.g243 314610 F M0004253C-DE01 UC2-NormColon 5977 457772 1561.L15.g243 314621 F M0004253C-DE01 UC2-NormColon 5977	5965	459974		F	M00042537C:D04	UC2-NormColon
5968 457846 1561.GI1_g243_314559 F M00042527B:B01 UC2-NormColon 5969 15296 15296 15296 15296 UC2-NormColon 5970 446578 1561.CI2_g243_314571 F M00042530C:H02 UC2-NormColon 5971 46308 1561.FI2_g243_314577 F M00042530D:A02 UC2-NormColon 5972 464287 1561.L12_g243_314577 F M00042530D:A02 UC2-NormColon 5973 447805 1561.L14_g243_314607 F M00042527C:A10 UC2-NormColon 5974 446188 1561.G14_g243_314610 F M00042527C:A10 UC2-NormColon 5975 476060 1561.L14_g243_314610 F M00042537D:F09 UC2-NormColon 5976 476060 1561.L14_g243_314610 F M00042327A:B04 UC2-NormColon 5977 457772 1561.R15_g243_314621 F M00042324D:B10 UC2-NormColon 5978 446731 1561.E15_g243_314621 F M00042360 UC2-NormColon 5978	5966	1454	1561.N10.gz43_314550	F	M00042539D:A04	UC2-NormColon
15296	5967	464510	1561.D11.gz43_314556	F	M00042522D:A08	UC2-NormColon
5970 446578 1561.C12.gs43_314571 F M00042345D-D04 UC2-NormColon 5971 463086 1561.F12.gs43_314577 F M00042525C:H07 UC2-NormColon 5972 464287 1561.H12.gs43_314577 F M00042530D:A02 UC2-NormColon 5973 447805 1561.H12.gs43_314601 F M0004254ID:G11 UC2-NormColon 5974 446188 1561.G14.gs43_314609 F M00042530D:F09 UC2-NormColon 5976 456565 1561.H4.gs43_314610 F M00042530D:F09 UC2-NormColon 5977 457772 1561.A15.gs43_314610 F M00042532A:H03 UC2-NormColon 5978 446733 1561.H15.gs43_314621 F M0004253CB:D01 UC2-NormColon 5979 446673 1561.F15.gs43_314622 F M0004253GB:G08 UC2-NormColon 5980 462779 1561.L15.gs43_314623 F M0004253GB:G08 UC2-NormColon 5981 446289 1561.L16.gs43_314643 F M0004253GB:G08 UC2-NormColon	5968	457846	1561.G11.gz43_314559	F	M00042527B:B01	UC2-NormColon
3971 463086 1561,F12,g243_314574 F M00042525C:H07 UC2-NormColon 5972 464287 1561,L12,g243_314577 F M00042530D:A02 UC2-NormColon 5973 447805 1561,L14,g243_314607 F M00042530D:A02 UC2-NormColon 5974 446188 1561,G14,g243_314607 F M00042527C:A10 UC2-NormColon 5976 476060 1561,L14,g243_314610 F M00042532A:H03 UC2-NormColon 5977 457772 1561,A15,g243_314610 F M00042352A:H03 UC2-NormColon 5978 446733 1561,E15,g243_314621 F M00042342A:B04 UC2-NormColon 5979 446673 1561,E15,g243_314621 F M00042350B:G08 UC2-NormColon 5980 462779 1561,L15,g243_314635 F M0004236B:B08 UC2-NormColon 5981 446289 1561,L16,g243_314643 F M0004253G:C06 UC2-NormColon 5981 445699 1561,L16,g243_314644 F M0004253G:C06 UC2-NormColon	5969	15296	1561.I11.gz43_314561	F	M00042530C:H02	UC2-NormColon
5972 464287 1561.112.g243_314577 F M00042530D.A02 UC2-NormColon 5973 447805 1561.A14.g243_314601 F M00042341D.G11 UC2-NormColon 5974 446188 1561.G14.g243_314607 F M00042527C.A10 UC2-NormColon 5975 468565 1561.114.g243_314609 F M00042530D.F09 UC2-NormColon 5976 477020 1561.A14.g243_314610 F M00042532A.H03 UC2-NormColon 5977 4577772 1561.A15.g243_314617 F M00042524C-C12 UC2-NormColon 5978 446383 1561.E15.g243_314622 F M0004253D.E01 UC2-NormColon 5980 462779 1561.L15.g243_314622 F M0004253D.E01 UC2-NormColon 5981 446289 1561.C16.g243_314643 F M0004253H.B08 UC2-NormColon 5981 445288 1561.N16.g243_314644 F M0004253GB.G08 UC2-NormColon 5984 45288 1561.N17.g243_314650 F M0004254D.B03 UC2-NormColon	5970	446578	1561.C12.gz43_314571	F	M00042345D:D04	UC2-NormColon
5973 447805 1561.A14 gz43 314601 F M0004234ID.G11 UC2-NormColon 5974 446188 1561.G14 gz43 314607 F M0004232TC:A10 UC2-NormColon 5975 468565 1561.H14 gz43 314609 F M0004253DF.P09 UC2-NormColon 5976 470600 1561.J14 gz43 314610 F M0004233ADF.P09 UC2-NormColon 5977 445772 1561.A15 gz43 314617 F M0004232AC:B04 UC2-NormColon 5978 446383 1561.B15 gz43 314621 F M0004252AC:C12 UC2-NormColon 5979 446673 1561.F15 gz43 314622 F M0004252G-C10 UC2-NormColon 5981 446289 1561.L16 gz43 314643 F M0004253GB:G08 UC2-NormColon 5982 458146 1561.K16 gz43 314643 F M0004253GB:G08 UC2-NormColon 5984 452888 1561.N16 gz43 314644 F M0004254AB:B08 UC2-NormColon 5984 452888 1561.N17 gz43 314650 F M0004234AB:P02 UC2-NormColon	5971	463086	1561.F12.gz43_314574	F	M00042525C:H07	UC2-NormColon
3974 446188 1561.G14.g243 314607 F M00042527C:A10 UC2-NormColon 3975 468365 1561.L14.g243 314609 F M00042530D:F09 UC2-NormColon 3976 47060 1561.114.g243 314610 F M00042530D:F09 UC2-NormColon 3977 47772 1561.A15.g243 314617 F M00042542A:B04 UC2-NormColon 3978 446373 1561.E15.g243 314621 F M0004253C:C12 UC2-NormColon 5979 446673 1561.E15.g243 314622 F M0004253G:B0 UC2-NormColon 5980 462779 1561.L15.g243 314635 F M0004253G:B0 UC2-NormColon 5981 446289 1561.L16.g243 314643 F M0004253G:D60 UC2-NormColon 5982 481146 1561.L16.g243 314644 F M0004254B:B03 UC2-NormColon 5984 45288 1561.N16.g243 314650 F M0004254B:B03 UC2-NormColon 5987	5972	464287	1561.I12.gz43_314577	F	M00042530D:A02	UC2-NormColon
5975 468565 1561.114.gz43_314609 F M00042530D.F99 UC2-NormColon 5976 470600 1561.114.gz43_314610 F M00042532A.FH03 UC2-NormColon 5977 457772 1561.A15.gz43_314617 F M00042532A.FH03 UC2-NormColon 5978 446383 1561.E15.gz43_314621 F M00042524C:C12 UC2-NormColon 5979 446673 1561.F15.gz43_314622 F M0004253ED:E01 UC2-NormColon 5980 462779 1561.L15.gz43_314632 F M0004253GB:G08_UC2-NormColon 5981 446289 1561.C16.gz43_314643 F M0004253HB:08 UC2-NormColon 5981 445288 1561.N16.gz43_314644 F M0004253GB:G08_UC2-NormColon 5984 45288 1561.N16.gz43_314646 F M0004254DB:B03 UC2-NormColon 5984 45288 1561.N17.gz43_314650 F M0004234B:F02 UC2-NormColon 5984 45288 1561.R17.gz43_314650 F M0004234B:F02 UC2-NormColon 5984 4599	5973	447805	1561.A14.gz43_314601	F	M00042341D:G11	UC2-NormColon
5976 470600 1561,114,gz43 314610 F M00042532A;H03 UC2-NormColon 5977 457772 1561,A15,gz43 314617 F M00042342A;B04 UC2-NormColon 5978 44633 1561,B15,gz43 314621 F M00042524C;C12 UC2-NormColon 5979 446673 1561,F15,gz43 314622 F M0004253GB;G08 UC2-NormColon 5980 466279 1561,L15,gz43 314628 F M0004253GB;G08 UC2-NormColon 5981 446289 1561,C16,gz43 314635 F M0004253GB;G08 UC2-NormColon 5982 458146 1561,K16,gz43 314643 F M0004253GB;008 UC2-NormColon 5984 45288 1561,L16,gz43 314646 F M00042540B;B03 UC2-NormColon 5985 461327 1561,B17,gz43 314650 F M00042544B;F02 UC2-NormColon 5987 467901 1561,K17,gz43 314654 F M00042540C;A12 UC2-NormColon 5988<	5974	446188	1561.G14.gz43_314607	F	M00042527C:A10	UC2-NormColon
5977 457772 1561.A15_gz43_314617 F M00042342A:B04 UC2-NormColon 5978 446383 1561.B15_gz43_314621 F M00042524C:C12 UC2-NormColon 5980 446673 1561.B15_gz43_314622 F M00042523D:E01 UC2-NormColon 5980 462779 1561.L15_gz43_314628 F M0004253G:B08 UC2-NormColon 5981 446289 1561.L16_gz43_314635 F M0004253G:B08 UC2-NormColon 5982 458146 1561.K16_gz43_314643 F M0004253G:B08 UC2-NormColon 5983 447690 1561.L16_gz43_314644 F M00042346:B03 UC2-NormColon 5984 452888 1561.N16_gz43_314650 F M0004234B:P02 UC2-NormColon 5985 461327 1561.B17_gz43_314650 F M0004254D:B03 UC2-NormColon 5987 467901 1561.F17_gz43_314654 F M0004254D:B03 UC2-NormColon 5989 45218 1561.N17_gz43_314665 F M0004234B:D06 UC2-NormColon <	5975	468565	1561.I14.gz43_314609	F	M00042530D:F09	UC2-NormColon
5978 446383 1561.E15 gz43 314621 F M00042524C:C12 UC2-NormColon 5979 446673 1561.F15 gz43 314622 F M00042532D:E01 UC2-NormColon 5980 46279 1561.L15 gz43 314635 F M00042531B:B08 UC2-NormColon 5981 446289 1561.C16 gz43 314643 F M00042531A:B08 UC2-NormColon 5982 445288 1561.N16 gz43 314644 F M00042534B:B08 UC2-NormColon 5984 452888 1561.N16 gz43 314644 F M00042534B:B03 UC2-NormColon 5985 461327 1561.B17 gz43 314650 F M0004234B:F02 UC2-NormColon 5986 456988 1561.R17 gz43 314653 F M0004234B:F02 UC2-NormColon 5987 467901 1561.F17 gz43 314653 F M0004234B:B00 UC2-NormColon 5989 457218 1561.N17 gz43 314659 F M0004234B:B00 UC2-NormColon 5989 457218 1561.N17 gz43 314650 F M00042342CH03 UC2-NormColon	5976	470600	1561.J14.gz43_314610	F	M00042532A:H03	UC2-NormColon
5979 446673 1561,F15,g243 314622 F M00042525B-E01 UC2-NormColon 5980 462779 1561,L15,g243 314628 F M00042526B-G08 UC2-NormColon 5981 446289 1561,L16,g243 314643 F M00042521A-E10 UC2-NormColon 5982 458146 1561,L16,g243 314643 F M00042534B-B08 UC2-NormColon 5984 452888 1561,L16,g243 314646 F M00042540B-B03 UC2-NormColon 5985 461327 1561,B17,g243 314650 F M00042344B-F02 UC2-NormColon 5986 456958 1561,B17,g243 314654 F M0004234AB-F02 UC2-NormColon 5987 467901 1561,K17,g243 314654 F M0004234AB-B03 UC2-NormColon 5989 457218 1561,K17,g243 314654 F M0004234C-A10 UC2-NormColon 5989 457218 1561,K17,g243 314665 F M00042340-A12 UC2-NormColon 5990<	5977	457772	1561.A15.gz43_314617	F	M00042342A:B04	UC2-NormColon
5980 462779 1561.L15.gz43_314628 F M0004253GB:G08 UC2-NormColon 5981 446289 1561.C16.gz43_314633 F M0004253LE.B0 UC2-NormColon 5982 4458146 1561.K16.gz43_314644 F M0004253GB:G08 UC2-NormColon 5983 447690 1561.L16.gz43_314644 F M0004236C:A06 UC2-NormColon 5984 452888 1561.N16.gz43_314650 F M0004234B:F02 UC2-NormColon 5985 461327 1561.B17.gz43_314650 F M0004234B:F02 UC2-NormColon 5986 456958 1561.E17.gz43_314653 F M0004234B:F02 UC2-NormColon 5987 467901 1561.F17.gz43_314659 F M0004234B:D6 UC2-NormColon 5988 457218 1561.N17.gz43_314669 F M0004234C:A12 UC2-NormColon 5990 46308 1561.L18.gz43_314669 F M0004234C:H03 UC2-NormColon 5991 446341 1561.L18.gz43_314669 F M00042342A:C11 UC2-NormColon <	5978	446383	1561.E15.gz43_314621	F	M00042524C:C12	UC2-NormColon
5981 446289 1561.C16.gz43 314635 F M00042521A:E10 UC2-NormColon 5982 458146 1561.K16.gz43 314643 F M00042534B:B08 UC2-NormColon 5983 447690 1561.L16.gz43 314646 F M00042530B:B03 UC2-NormColon 5984 452888 1561.N16.gz43 314646 F M00042540B:B03 UC2-NormColon 5985 461327 1561.B17.gz43 314650 F M00042540B:B03 UC2-NormColon 5987 467901 1561.F17.gz43 314653 F M00042534B:D0 UC2-NormColon 5988 455988 1561.F17.gz43 314654 F M00042536A:E10 UC2-NormColon 5989 457218 1561.N17.gz43 314650 F M00042536A:E10 UC2-NormColon 5990 463008 1561.A18.gz43 314665 F M00042342C:H03 UC2-NormColon 5991 446341 1561.L18.gz43 314667 F M00042536C:B03 UC2-NormColon 5992	5979	446673	1561.F15.gz43_314622	F	M00042525D:E01	UC2-NormColon
5982 458146 1561.K16.gz43_314643 F M00042534B:B08 UC2-NormColon 5983 447690 1561.L16.gz43_314644 F M00042536C:A06 UC2-NormColon 5984 452888 1561.N16.gz43_314646 F M00042540B:B03_UC2-NormColon 5985 461327 1561.B17.gz43_314650 F M00042344B:F02 UC2-NormColon 5986 456958 1561.E17.gz43_314653 F M00042324D:B03_UC2-NormColon UC2-NormColon 5987 467901 1561.E17.gz43_314654 F M00042534B:D06 UC2-NormColon 5988 459981 1561.K17.gz43_314665 F M00042534B:D06 UC2-NormColon 5980 467308 1561.N17.gz43_314665 F M0004254C:A12 UC2-NormColon 5990 463008 1561.L18.gz43_314667 F M00042534C:H03_UC2-NormColon 5991 446341 1561.L18.gz43_314676 F M00042534C:H03_UC2-NormColon 5992 46784 1561.B19.gz43_314680 F M00042542A:C11 UC2-NormColon 5993 44709	5980	462779	1561.L15.gz43_314628	F	M00042536B:G08	UC2-NormColon
5983 447690 1561.L16.gz43_314644 F M00042536C:A06 UC2-NormColon 5984 45288 1561.N16.gz43_314664 F M00042540B:B03 UC2-NormColon 5985 461327 1561.B17.gz43_314653 F M0004254D:A10 UC2-NormColon 5986 456958 1561.B17.gz43_314653 F M0004252AD:A10 UC2-NormColon 5987 467901 1561.F17.gz43_314659 F M0004252AD:A10 UC2-NormColon 5988 459981 1561.K17.gz43_314659 F M0004252AD:D0 UC2-NormColon 5989 457218 1561.N17.gz43_314665 F M0004254CA:L10 UC2-NormColon 5990 463008 1561.A18.gz43_314665 F M00042342C:H03 UC2-NormColon 5991 446341 1561.L18.gz43_314676 F M00042342A:C11 UC2-NormColon 5992 446389 1561.L18.gz43_314680 F M00042342A:C11 UC2-NormColon 5994 446784 1561.B19.gz43_314680 F M00042342D:H11 UC2-NormColon	5981	446289	1561.C16.gz43_314635	F	M00042521A:E10	UC2-NormColon
5984 452888 1561.N16.gz43 314646 F M00042540B:B03 UC2-NormColon 5985 461327 5151.B17.gz43 314650 F M0004234B:F02 UC2-NormColon 5986 45698 1561.B17.gz43 314653 F M00042324B:A10 UC2-NormColon 5987 467901 1561.F17.gz43 314654 F M00042536A:E10 UC2-NormColon 5988 459981 1561.K17.gz43 314650 F M00042536A:E10 UC2-NormColon 5990 463008 1561.A18.gz43 314665 F M00042342C:H03 UC2-NormColon 5991 446341 1561.L18.gz43 314676 F M00042342C:H03 UC2-NormColon 5992 446398 1561.D18.gz43 314676 F M00042536C:B03 UC2-NormColon 5993 447096 1561.P18.gz43 314680 F M00042543B:H12 UC2-NormColon 5994 46784 1561.B19.gz43 314680 F M00042254B:G11 UC2-NormColon 5995 </td <td>5982</td> <td>458146</td> <td>1561.K16.gz43_314643</td> <td>F</td> <td>M00042534B:B08</td> <td>UC2-NormColon</td>	5982	458146	1561.K16.gz43_314643	F	M00042534B:B08	UC2-NormColon
5985 461327 1561.B17 gz43_314650 F M00042344B:F02 UC2-NormColon 5986 456958 1561.B17 gz43_314653 F M00042324D:A10 UC2-NormColon 5987 467901 1561.F17 gz43_314654 F M00042326D:A10 UC2-NormColon 5988 459981 1561.K17 gz43_314659 F M0004234B:D06 UC2-NormColon 5980 457128 1561.N17 gz43_314665 F M00042340:CA12 UC2-NormColon 5990 463008 1561.A18 gz43_314665 F M00042342C:H03 UC2-NormColon 5991 446341 1561.L18 gz43_314676 F M00042342A:CH03 UC2-NormColon 5992 446381 1561.O18 gz43_314679 F M00042342A:CH1 UC2-NormColon 5993 447096 1561.P18 gz43_314680 F M00042343B:H12 UC2-NormColon 5995 468767 1561.B19 gz43_314680 F M0004234B:H12 UC2-NormColon 5995 467884 1561.H19 gz43_314680 F M00042320 UC2-NormColon	5983	447690	1561.L16.gz43_314644	F	M00042536C:A06	UC2-NormColon
5986 456958 1561.E17.gz43_314653 F M00042524D:A10 UC2-NormColon 5987 467901 1561.F17.gz43_314654 F M00042526A:E10 UC2-NormColon 5988 459981 1561.K17.gz43_314659 F M00042534B:D06 UC2-NormColon 5989 457218 1561.K17.gz43_314662 F M00042540C:A12 UC2-NormColon 5990 463008 1561.A18.gz43_314665 F M00042342C:H03 UC2-NormColon 5991 446341 1561.L18.gz43_314676 F M00042342A:C11 UC2-NormColon 5992 446389 1561.D18.gz43_314680 F M00042342A:C11 UC2-NormColon 5993 447096 1561.P18.gz43_314680 F M00042343HH12 UC2-NormColon 5994 446984 1561.B19.gz43_314680 F M00042342HH12 UC2-NormColon 5995 467876 1561.F19.gz43_314686 F M000423263-F11 UC2-NormColon 5997 495881 1561.A20.gz43_314697 F M0004232D:B03 UC2-NormColon	5984	452888	1561.N16.gz43_314646	F	M00042540B:B03	UC2-NormColon
5987 467901 1561.F17.gz43_314654 F M00042526A:E10 UC2-NormColon 5988 457981 1561.K17.gz43_314659 F M00042534B:D06 UC2-NormColon 5989 457218 1561.N17.gz43_314665 F M00042340C:A12 UC2-NormColon 5990 463008 1561.A18.gz43_314665 F M00042342C:H03 UC2-NormColon 5991 446341 1561.L18.gz43_314676 F M00042342C:H03 UC2-NormColon 5992 446398 1561.D18.gz43_314676 F M00042534C:B03 UC2-NormColon 5993 447096 1561.P18.gz43_314680 F M00042543A:B112 UC2-NormColon 5994 446384 1561.B19.gz43_314680 F M0004234B:H12 UC2-NormColon 5995 463767 1561.F19.gz43_314680 F M00042329B:E03 UC2-NormColon 5995 467884 1561.H19.gz43_314680 F M00042329B:E03 UC2-NormColon 5997 459881 1561.A20.gz43_314697 F M00042342D:D03 UC2-NormColon	5985	461327	1561.B17.gz43_314650	F	M00042344B:F02	UC2-NormColon
5988 459981 1561.K17_g243_314659 F M00042534B:D06 UC2-NormColon 5989 457218 1561.N17_g243_314662 F M00042540C:A12 UC2-NormColon 5990 46308 1561.A18_g243_314665 F M00042342C:H03 UC2-NormColon 5991 446341 1561.L18_g243_314676 F M000425342A:C11 UC2-NormColon 5992 446389 1561.D18_g243_314679 F M00042542A:C11 UC2-NormColon 5993 447096 1561.P18_g243_314686 F M00042543B:H12 UC2-NormColon 5995 468767 1561.F19_g243_314686 F M0004254B:H12 UC2-NormColon 5996 467884 1561.H19_g243_314686 F M00042529B:E03 UC2-NormColon 5997 459881 1561.A20_g243_314697 F M0004232D:D03 UC2-NormColon 5998 447027 1561.D20_g243_314700 F M0004232B:H01 UC2-NormColon 6000 446304 1561.O21_g243_314771 F M0004252B:B01 UC2-NormColon	5986	456958	1561.E17.gz43_314653	F	M00042524D:A10	UC2-NormColon
5989 457218 1561.N17.gz43_314662 F M00042540C:A12 UC2-NormColon 5990 463008 1561.A18.gz43_314665 F M00042342C:H03 UC2-NormColon 5991 446381 1561.L18.gz43_314679 F M00042542A:C11 UC2-NormColon 5992 446389 1561.D18.gz43_314689 F M00042542A:C11 UC2-NormColon 5993 447096 1561.P18.gz43_314680 F M0004234B:H12 UC2-NormColon 5994 446984 1561.P18.gz43_314680 F M0004234B:H12 UC2-NormColon 5995 468767 1561.F19.gz43_314686 F M00042526A:F11 UC2-NormColon 5996 467884 1561.H19.gz43_314687 F M00042529B:E03 UC2-NormColon 5997 449881 1561.A20.gz43_314697 F M00042529B:E03 UC2-NormColon 5998 447072 1561.D20.gz43_314702 F M00042526B:C12 UC2-NormColon 5999 447380 1561.O21.gz43_314773 F M0004252B:B01 UC2-NormColon	5987	467901	1561.F17.gz43_314654	F	M00042526A:E10	UC2-NormColon
5990 463008 1561.A18_gz43_314665 F M00042342C:H03 UC2-NormColon 5991 446341 1561.L18_gz43_314676 F M00042536C:B03 UC2-NormColon 5992 446389 1561.O18_gz43_314676 F M00042542A:C11 UC2-NormColon 5993 447096 1561.P18_gz43_314680 F M00042543B:H12 UC2-NormColon 5994 446984 1561.B19_gz43_314680 F M0004234B:H12 UC2-NormColon 5995 468767 1561.F19_gz43_314686 F M00042326A:F11 UC2-NormColon 5996 467884 1561.H19_gz43_314686 F M00042529B:E03 UC2-NormColon 5997 45981 1561.A20_gz43_314697 F M00042342D:D03 UC2-NormColon 5998 47027 1561.D20_gz43_314700 F M0004252B:H01 UC2-NormColon 5999 447380 1561.C21_gz43_314720 F M0004252B:B01 UC2-NormColon 6001 447072 1561.C22_gz43_314731 F M0004252B:DG9 UC2-NormColon	5988	459981	1561.K17.gz43_314659	F	M00042534B:D06	UC2-NormColon
5991 446341 1561.L18_gz43_314676 F M00042536C:B03 UC2-NormColon 5992 446389 1561.O18_gz43_314679 F M00042542A:C11 UC2-NormColon 5993 447096 1561.P18_gz43_314680 F M00042243B:H12 UC2-NormColon 5994 44684 1561.B19_gz43_314686 F M0004224B:H12 UC2-NormColon 5995 468767 1561.F19_gz43_314686 F M00042256A:F11 UC2-NormColon 5996 467884 1561.H19_gz43_314688 F M00042529E:B3 UC2-NormColon 5997 459881 1561.A20_gz43_314697 F M00042529B:B3 UC2-NormColon 5998 447027 1561.A20_gz43_314702 F M0004252B:B11 UC2-NormColon 6000 446304 1561.O21_gz43_314727 F M0004252B:B01 UC2-NormColon 6001 447072 1561.O21_gz43_314731 F M0004252B:DG UC2-NormColon 6002 379335 1561.E22_gz43_314737 F M0004253IB:D12 UC2-NormColon	5989	457218	1561.N17.gz43_314662	F	M00042540C:A12	UC2-NormColon
5992 446389 1561.018.gz43_314679 F M00042542A:C11 UC2-NormColon 5993 447096 1561.P18.gz43_314680 F M00042543B:H12 UC2-NormColon 5994 446984 1561.B19.gz43_314680 F M0004234B:H12 UC2-NormColon 5995 468767 1561.F19.gz43_314686 F M0004232A:B:H13 UC2-NormColon 5996 467884 1561.H19.gz43_314687 F M0004232B:B03 UC2-NormColon 5997 459881 1561.A20.gz43_314697 F M0004252B:B03 UC2-NormColon 5998 447027 1561.D20.gz43_314700 F M0004252B:C12 UC2-NormColon 5999 447380 1561.F20.gz43_314720 F M0004252B:D1 UC2-NormColon 6000 446304 1561.O21.gz43_314727 F M0004252B:D1 UC2-NormColon 6001 447072 1561.C22.gz43_314731 F M0004252B:D6 UC2-NormColon 6002 379335 1561.U22.gz43_314731 F M0004252ID:G09 UC2-NormColon	5990	463008	1561.A18.gz43_314665	F	M00042342C:H03	UC2-NormColon
5993 447096 1561.P18.g243_314680 F M00042543B:H12 UC2-NormColon 5994 446984 1561.B19.g243_314682 F M0004234B:H12 UC2-NormColon 5995 46876 1561.F19.g243_314688 F M00042526A:F11 UC2-NormColon 5996 467884 1561.H19.g243_314688 F M00042529B:E03 UC2-NormColon 5997 459881 1561.A20.g243_314697 F M00042342D:D03 UC2-NormColon 5998 447072 1561.D20.g243_314700 F M0004252B:B01 UC2-NormColon 5999 447300 1561.C21.g243_314727 F M00042542B:B01 UC2-NormColon 6000 446304 1561.C21.g243_314737 F M00042521D:G09 UC2-NormColon 6001 447072 1561.C22.g243_314737 F M00042531B:D12 UC2-NormColon 6002 379335 1561.U22.g243_314737 F M00042531B:D12 UC2-NormColon	5991	446341	1561.L18.gz43_314676	F	M00042536C:B03	UC2-NormColon
5994 446984 1561.B19 gr43 314682 F M00042344BH12 UC2-NormColon 5995 46787 1561.F19 gr43 314686 F M00042356A:F11 UC2-NormColon 5996 46784 1561.H19 gr43 314688 F M00042329B:E03 UC2-NormColon 5997 459881 1561.A20 gr43 314697 F M00042342D:D03 UC2-NormColon 5998 447027 1561.D20 gr43 314700 F M00042342B:D10 UC2-NormColon 6000 446304 1561.O21 gr43 314727 F M00042542B:B01 UC2-NormColon 6001 447072 1561.O21 gr43 314737 F M00042521D:G09 UC2-NormColon 6002 379335 1561.U22 gr43 314737 F M00042531B:D12 UC2-NormColon	5992	446389	1561.O18.gz43_314679	F	M00042542A:C11	UC2-NormColon
5995 468767 1561.F19 g-843 314686 F M00042526A:F11 UC2-NormColon 5996 467884 1561.H19 g-843 314688 F M00042529B:E03 UC2-NormColon 5997 459881 1561.A20 g-843 314697 F M00042329B:E03 UC2-NormColon 5998 447027 1561.D20 g-843 314700 F M00042523B:H01 UC2-NormColon 5999 447380 1561.F20 g-843 314720 F M00042526B:C12 UC2-NormColon 6000 446304 1561.O21 g-843 314727 F M0004252B:B01 UC2-NormColon 6001 447072 1561.C22 g-843 314731 F M0004252B:B01 UC2-NormColon 6002 379335 1561.U22 g-843 314731 F M0004253B:D10 UC2-NormColon 6002 379335 1561.U22 g-843 314737 F M0004253B:D10 UC2-NormColon	5993	447096	1561.P18.gz43_314680	F	M00042543B:H12	UC2-NormColon
5996 467884 1561.H19 zp43 314688 F M00042529B:E03 UC2-NormColon 5997 459881 1561.A20 zp43 314697 F M00042342D:D03 UC2-NormColon 5998 447072 1561.D20 zp43 314700 F M00042523B:H01 UC2-NormColon 5999 447380 1561.F20 zp43 314702 F M00042526B:C12 UC2-NormColon 6000 446304 1561.O21 zp43 314727 F M00042542B:B01 UC2-NormColon 6001 447072 1561.C22 zp43 314731 F M00042521D:G09 UC2-NormColon 6002 379335 1561.U22 zp43 314737 F M00042531B:D12 UC2-NormColon	5994	446984	1561.B19.gz43_314682	F	M00042344B:H12	UC2-NormColon
5997 459881 1561.A20.gz43_314697 F M00042342D:D03 UC2-NormColon 5998 447027 1561.D20.gz43_314700 F M00042523B:H01 UC2-NormColon 5999 447380 1561.F20.gz43_314702 F M00042526B:C12 UC2-NormColon 6000 446304 1561.O21.gz43_314727 F M0004252B:D61 UC2-NormColon 6001 447072 1561.C22.gz43_314731 F M00042521D:G09 UC2-NormColon 6002 379335 1561.I22.gz43_314737 F M00042531B:D12 UC2-NormColon	5995	468767	1561.F19.gz43_314686	F	M00042526A:F11	UC2-NormColon
5998 447027 1561.D20.gz43_314700 F M00042523B:H01 UC2-NormColon 5999 447380 1561.F20.gz43_314702 F M00042526B:C12 UC2-NormColon 6000 446304 1561.O21.gz43_314727 F M0004252BB:I UC2-NormColon 6001 447072 1561.C22.gz43_314731 F M0004252ID.G09 UC2-NormColon 6002 379335 1561.I22.gz43_314737 F M0004253IB:D12 UC2-NormColon	5996	467884	1561.H19.gz43_314688	F	M00042529B:E03	UC2-NormColon
5999 447380 1561.F20.gz43_314702 F M00042526B:C12 UC2-NormColon 6000 446304 1561.021.gz43_314727 F M00042542B:B01 UC2-NormColon 6001 447072 1561.C22.gz43_314731 F M00042521D:G09 UC2-NormColon 6002 379335 1561.U22.gz43_314737 F M00042531B:D12 UC2-NormColon	5997	459881	1561.A20.gz43_314697	F	M00042342D:D03	UC2-NormColon
6000 446304 1561.021.gz/43 314727 F M00042542B:B01 UC2-NormColon 6001 447072 1561.C22.gz/43 314731 F M00042521D:G09 UC2-NormColon 6002 379335 1561.122.gz/43 314737 F M00042531B:D12 UC2-NormColon	5998	447027	1561.D20.gz43_314700	F	M00042523B:H01	UC2-NormColon
6001 447072 1561.C22.gz43_314731 F M00042521D:G09 UC2-NormColon 6002 379335 1561.I22.gz43_314737 F M00042531B:D12 UC2-NormColon	5999	447380	1561.F20.gz43_314702	F	M00042526B:C12	
6002 379335 1561.I22.gz43_314737 F M00042531B:D12 UC2-NormColon	6000	446304	1561.021.gz43_314727	F	M00042542B:B01	UC2-NormColon
The state of the s	6001	447072	1561.C22.gz43_314731	F	M00042521D:G09	UC2-NormColon
6002 229972 1561 122 -42 214740 E MODOMASSA ADT 1102 N C.1.	6002	379335	1561.I22.gz43_314737	F	M00042531B:D12	UC2-NormColon
0003 228873 1301,L22,g243_314740 F M00042537A:A07 UC2-NormColon	6003	228873	1561,L22.gz43_314740	F	M00042537A:A07	UC2-NormColon
6004 446663 1561.D23.gz43_314748 F M00042523C:E08 UC2-NormColon	6004	446663	1561.D23.gz43_314748	F	M00042523C:E08	UC2-NormColon
6005 464791 1561.F23.gz43_314750 F M00042526C:B12 UC2-NormColon	6005	464791	1561.F23.gz43_314750	F	M00042526C:B12	UC2-NormColon
6006 446595 1561.K23.gz43_314755 F M00042534D:D10 UC2-NormColon	6006	446595	1561.K23.gz43_314755	F	M00042534D:D10	UC2-NormColon

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Table 2

SEQ					
ID			ORIE		
NO	CLUSTER	SEQ NAME	NT	CLONE ID	LIBRARY
6007	446439	1561.023.gz43_314759	F	M00042542B:C11	UC2-NormColon
6008	458979	1561.A24.gz43_314761	F	M00042343A:C05	UC2-NormColon
6009	446620	1561.C24.gz43_314763	F	M00042522A:A05	UC2-NormColon
6010	452507	1561.F24.gz43_314766	F	M00042526D:A02	UC2-NormColon

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	-
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
1	AJ251957	Ciona intestinalis mRNA for nuclear lamin (lamin L2 gene)	2.2
2	AF133812	Strongylocentrotus pallidus isolate pR31 bindin gene, partial cds	0.88
3	AF051651	Homo sapiens squalene synthase gene, intron 3, 5' end	0.0001
4	NM_016701	Mus musculus nestin (Nes), mRNA	0.24
5	Y09000	R.norvegicus mRNA for dendrin	0.11
6	U67519	Methanococcus jannaschii section 61 of 150 of the complete genome	6.1
_		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
7	AK005889	library, clone: 1700012B08, full insert sequence	e-139
		Homo sapiens HCR (a-helix coiled-coil rod homologue) gene,	1
8	AB029343	complete cds	0.023
9	AB017505	Homo sapiens mRNA for Nori-2p, complete cds	0
		Oryza sativa microsatellite MRG6571 containing (TCTA)X6,	
10	AY024246	genomic sequence	6.3
		Acipenser fulvescens Af14 cytochrome b gene, partial cds, tRNA-Thr,	
		tRNA-Pro and partial D-loop, mitochondrial genes encoding	
11	U32309	mitochondrial products	6.3
12	XM_046070	Homo sapiens KIAA0853 protein (KIAA0853), mRNA	3E-48
13	AK001945	Homo sapiens cDNA FLJ11083 fis, clone PLACE1005232	2.2
14	AF111457	Meleagris gallopavo clone TUCA1141 microsatellite sequence	0.076
		Mus musculus adult male colon cDNA, RIKEN full-length enriched	
15	AK018499	library, clone:9030409O13, full insert sequence	١ ٥
		Drosophila melanogaster genomic scaffold 142000013385436,	
16	AE002648	complete sequence	2.2
		Homo sapiens (subclone 1_e5 from P1 H43) DNA sequence, complete	
17	AC001011	sequence	0.003
18	XM 039942	Homo sapiens Meis (mouse) homolog 3 (MEIS3), mRNA	0.75
19	AF099003	Caenorhabditis elegans cosmid Y59C2A	0.75
		Mus musculus protease-activated receptor 3 (PAR3) mRNA, complete	
20	U92972	cds	0.08
			0.00
21	AE003935	Xylella fastidiosa 9a5c, section 81 of 229 of the complete genome	0.067
		- 1, 1 that and a south of the south of the complete general	0,001
22	AB031009	Homo sapiens DNA, MHC class I CL region, 7.1 ancestral haplotype	0.019
23	X94207	H. sapiens TPR gene (1365bp)	4.8
24	AK025678	Homo sapiens cDNA: FLJ22025 fis, clone HEP08518	2.2
		Mus musculus 0 day neonate skin cDNA, RIKEN full-length enriched	
25	AK014552	library, clone:4632404O06, full insert sequence	e-176
26	XM 044013	Homo sapiens kinesin-like protein 2 (hklp2), mRNA	0
		Homo sapiens nucleolar protein family A, member 1 (H/ACA small	— <u> </u>
27	NM_032993	nucleolar RNPs) (NOLA1), transcript variant 2, mRNA	4E-50
28	NM_032315	Homo sapiens hypothetical protein MGC4399 (MGC4399), mRNA	0
		Mus musculus 6 days neonate skin cDNA, RIKEN full-length	
29	AK020701	enriched library, clone; A030009B12, full insert sequence	0,0000002
30	AK021717	Homo sapiens cDNA FLJ11655 fis, clone HEMBA1004554	0

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WC0214560 [fle //E /WO0214500 opc]

Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE Human DNA sequence from clone CTC-1516K22 on chromosome 6 31 AL512453 complete sequence [Homo sapiens] 0.00004 32 XM 005034 Homo sapiens hypothetical protein FLJ10511 (FLJ10511), mRNA 2E-22 Cricetulus griseus beta-1,4-galactosyltransferase 1 (beta4GalT-1) 33 AF318896 mRNA, complete cds 0.78 34 D29757 Human gene for cytochrome P-450 aromatase, exon 1 (brain specific) 3E-24 Ostreopsis ovata 5.8S rRNA gene, internal transcribed spacer 1 (ITS1) 35 AJ311520 and internal transcribed spacer 2 (ITS2), strain A1 0.79 XM 038521 Homo sapiens phosphatidylserine decarboxylase (PISD), mRNA 36 37 AK023043 Homo sapiens cDNA FLJ12981 fis, clone NT2RP2006454 3E-96 38 XM 045520 Homo sapiens KIAA1595 protein (KIAA1595), mRNA 4E-16 39 AB017505 Homo sapiens mRNA for Nori-2p, complete cds 0 Mus musculus adult male kidney cDNA, RIKEN full-length enriched AK002682 40 library, clone:0610027B03, full insert sequence e-138 NC 001712 41 Locusta migratoria mitochondrion, complete genome 0.009 Human DNA sequence from clone CTC-1516K22 on chromosome 6, 42 AL512453 complete sequence [Homo sapiens] 0.00001 Mus musculus adult male kidney cDNA, RIKEN full-length enriched 43 AK002682 library, clone:0610027B03, full insert sequence e-125 Homo sapiens KIAA0914 gene product (KIAA0914), mRNA 44 XM 039088 0.76 Schistosoma mansoni synaptobrevin-like protein gene, exon 2 and 45 U30291 partial cds 0.69 Homo sapiens mitochondrial ribosomal protein L42 (MRPL42), 46 XM 052543 1E-26 mRNA 47 AK026225 Homo sapiens cDNA: FLJ22572 fis, clone HSI02313 0.0008 Homo sapiens hypothetical gene supported by AK021969 48 XM 043007 (LOC92132), mRNA 1E-99 49 M18824 P.faciparum S antigen gene, complete cds 0.003 50 XM 009311 Homo sapiens zinc finger protein 304 (ZNF304), mRNA 0.003 AK026697 51 Homo sapiens cDNA: FLJ23044 fis, clone LNG02454 0 52 XM 035597 Homo sapiens hypothetical protein AF140225 (AF140225), mRNA 1E-98 53 U81366 Plasmid Rts1 killer protein and antidote protein genes, complete cds 0.26 Phaseolus vulgaris ATPase gamma subunit mRNA, nuclear gene 54 U28921 encoding mitochondrial protein, partial cds 0.27 55 AF305083 Homo sapiens alpha(1,3)-fucosyltransferase IV (FUTIV) gene, 3' UTR 0.031 56 D29757 Human gene for cytochrome P-450 aromatase, exon 1 (brain specific) 7E-20 Homo sapiens similar to APICAL ENDOSOMAL GLYCOPROTEIN 57 XM_038048 PRECURSOR (R. norvegicus) (LOC91372), mRNA 0.0004 AK026860 Homo sapiens cDNA: FLJ23207 fis, clone ADSE00968 8E-12

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Homo sapiens, Similar to hypothetical protein FLJ20561, clone	
59	BC010849	MGC:8851 IMAGE:3878776, mRNA, complete cds	0
60	XM_036092	Homo sapiens signal recognition particle 68kD (SRP68), mRNA	0
61	XM 003184	Homo sapiens zinc finger protein ANC_2H01 (LOC51193), mRNA	0
63	Z49806	B.taurus mRNA for adenyiyi cyclase type VII	6.8
		S. Maria S.	0.0
64	XM_048858	Homo sapiens hypothetical protein FLJ23384 (FLJ23384), mRNA	0
65	NM_032315	Homo sapiens hypothetical protein MGC4399 (MGC4399), mRNA	0
		Homo sapiens GSTT1 gene for glutathione S-transferase TT1,	
66	AB057594	complete cds	3E-69
67	XM 035597	Homo sapiens hypothetical protein AF140225 (AF140225), mRNA	2E-75
	_	, , , , , , , , , , , , , , , , , , ,	
68	XM 035597	Homo sapiens hypothetical protein AF140225 (AF140225), mRNA	1E-71
69	AB017505	Homo sapiens mRNA for Nori-2p, complete cds	0
	1115011505	Mus musculus adult male tongue cDNA, RIKEN full-length enriched	
70	AK010143	library, clone:2310074B19, full insert sequence	1.7
	111010113	Mus musculus adult male stomach cDNA, RIKEN full-length	1.7
71	AK008666	enriched library, clone:2210008A03, full insert sequence	e-173
/1	AKOOSOOO	entrened notary, crone.2210008A05, tun insert sequence	0-1/3
72	L18785	Plasmodium falciparum DNA polymerase alpha gene, complete cds	0.007
	110703	i iasinoulum iaiciparum Diva polymerase aipna gene, compiete cus	0.007
73	XM_003327	Homo sapiens hypothetical protein FLJ10858 (FLJ10858), mRNA	0
		Mus musculus adult male lung cDNA, RIKEN full-length enriched	
74	AK004786	library, clone:1200015F23, full insert sequence	2E-11
		E. histolytica plasmid genes for ribosomal RNA and hemolysins	
75	Z29969	HLY1, HLY5mc1 HLY5mc2 HLY4	0.028
76	AK024945	Homo sapiens cDNA: FLJ21292 fis, clone COL01969	0.020
77	Y15931	Homo sapiens CTNS gene, exon 10 and flanking intronic regions	0.78
		Homo sapiens, hypothetical protein FLJ10986, clone MGC:908	0.70
78	BC000610	IMAGE:3347108, mRNA, complete cds	0
79		Homo sapiens KIAA1244 protein (KIAA1244), mRNA	0.75
80	NC 001807	Human mitochondrion, complete genome	1E-36
- 50	1.5_001007	Arabidopsis thaliana unknown protein (F28C11.8) mRNA, complete	10-20
81	AF360266	cds	6.8
- 01	AL 500200	Homo sapiens, Similar to hypothetical protein FLJ10656, clone	0.0
82	BC000225	IMAGE:3350951, mRNA	0
- 02	130000223	INTERIOR, SOUSSE, MINISTER	
83	XM 034107	Homo sapiens hypothetical protein FLJ20758 (FLJ20758), mRNA	7E-32
84	AF305816	Homo sapiens PRO0633 mRNA, complete cds	0.67
	12130010	HIV-1 isolate 96RW34 from Rwanda, gag protein (gag) gene, partial	
85	AF196678	cds	2.1
86	AK024927	Homo sapiens cDNA: FLJ21274 fis, clone COL01781	6E-75
87	X77394	P.philodendra mitochondrial DNA repeat unit	0.23
- 01	*******	- Panoconata intochondra Dirix topoat unit	5,25

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
88	U41203	Human short tandem repeat polymorphism UM32, D17S1333	0.0001
89	AC015937	Homo sapiens chromosome 17, clone CTD-2542F12, complete sequence	0.65
90	AF245117	Mus musculus leukocyte cell-surface molecule (Ly9) gene, exon 1	0.74
91	XM_001412	Homo sapiens metal-regulatory transcription factor 1 (MTF1), mRNA	2E-50
92	AE006690	Sulfolobus solfataricus section 49 of 272 of the complete genome	0.26
93	AJ292466	Homo sapiens mRNA for WDR9 protein (WDR9 gene), form B	9E-44
94	NM_013983	Homo sapiens neuregulin 2 (NRG2), transcript variant 4, mRNA	0.27
95	J01390	Emericella nidulans mtDNA between h2/h5 and bh2/b2 junctions. genes for ATPase subunit 6, cytochrome oxidase subunit 3, seven. unidentified proteins, twentyfour tRNA's and L-rRNA	2.2
96	AP000384	Arabidopsis thaliana genomic DNA, chromosome 3, P1 clone:MCE21	0.22
97	AE001397	Plasmodium falciparum chromosome 2, section 34 of 73 of the complete sequence	2.4
98	AE000192	Escherichia coli K12 MG1655 section 82 of 400 of the complete genome	0.003
99	AF316828	Schistosoma mansoni zinc finger protein SmZF1 gene, complete cds	1.7
100	BC007942	Homo sapiens, nucleolar autoantigen (55kD) similar to rat synaptonemal complex protein, clone MGC:14267 IMAGE:4130726, mRNA, complete cds Homo sapiens nucleoside diphsophate kinase A (nm23-H1) gene	2E-59
101_	L35301	fragment	0.003
102	AF114927 AF299248	Saccharomyces pastorianus CBS1538 small subunit ribosomal RNA gene, mitochondrial gene for mitochondrial RNA, complete sequence Drosophila melanogaster talin mRNA, complete cds	0.27
103	AF299248		1,8
104	BC003191	Homo sapiens, GL004 protein, clone MGC:895 IMAGE:3502929, mRNA, complete cds	2E-10
105	U29926	Human AMP deaminase (AMPD3) gene, exon 14 and 15, and complete cds	0.06
106	BC003191	Homo sapiens, GL004 protein, clone MGC:895 IMAGE:3502929, mRNA, complete cds	0
107	AK022127	Homo sapiens cDNA FLJ12065 fis, clone HEMBB1002249	0,0008
108	D16360	Human DNA for plasma glutathione peroxidase, exon 1	0.091
109	AK014138	Mus musculus 13 days embryo head cDNA, RIKEN full-length enriched library, clone:3110038A11, full insert sequence	6
110	XM_006914	Homo sapiens cold shock domain protein A (CSDA), mRNA	0.079
111	U31283	Mesotaenium caldariorum clone mesphyla phytochrome gene, partial cds, exons 4 through 10	0.56
112	XM_049935	Homo sapiens hypothetical protein FLJ14950 (FLJ14950), mRNA	0

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
113	AE001329	Chlamydia trachomatis section 56 of 87 of the complete genome	0.75
114	XM_037830	Homo sapiens OLF-1/EBF associated zinc finger gene (KIAA0760), mRNA	0.25
115	AE006368	Lactococcus lactis subsp. lactis IL1403 section 130 of 218 of the complete genome	0.0003
116	XM_049935	Homo sapiens hypothetical protein FLJ14950 (FLJ14950), mRNA	0
117	XM_032533	Homo sapiens hypothetical protein FLJ12787 (FLJ12787), mRNA	0
		Homo sapiens partial mRNA for putative protein kinase WNK4	
118	AJ309861	(PRKWNK4 gene)	9E-76
l		Homo sapiens, KIAA0255 gene product, clone IMAGE:3507918,	
119	BC005173	mRNA	2E-38
120	AF068758	Drosophila melanogaster SIR2 (Sir2) mRNA, complete cds	0.083
		Mus musculus 6 days neonate head cDNA, RIKEN full-length	
121	AK017307	enriched library, clone:5430414B19, full insert sequence	0,003
122	XM_036208	Homo sapiens sphingosine kinase 1 (SPHK1), mRNA	2.1
123	XM_039960	Homo sapiens PC3-96 protein (PC3-96), mRNA	0
124	XM_039306	Homo sapiens hypothetical protein FLJ20400 (FLJ20400), mRNA	0,08
125	XM_049935	Homo sapiens hypothetical protein FLJ14950 (FLJ14950), mRNA	0
126	AE006080	Pasteurella multocida PM70 section 47 of 204 of the complete genome	2.1
127	XM_009518	Homo sapiens WNT1 inducible signaling pathway protein 2 (WISP2), mRNA	0.16
128	U36263	Homo sapiens beta-prime-adaptin (AP1B1) gene, exon 15	0.022
129	AF267996	Homo sapiens muscle-specific protein (C4orf5) gene, intron 3	0.34
130	AF131859	Homo sapiens clone 24923 mRNA sequence	1.8
131	L48479	Homo sapiens (subclone 6_h1 from P1 H21) DNA sequence	0.01
132	XM_018317	Homo sapiens ubiquitination factor E4B (homologous to yeast UFD2) (UBE4B), mRNA	1.6
133	XM_008267	Homo sapiens solute carrier family 16 (monocarboxylic acid transporters), member 6 (SLC16A6), mRNA	0,23
134	AF162923	Beet soil-borne virus RNA3 isolate 6b 22K protein gene, partial cds	6.6
135	NM_010053	Mus musculus distal-less homeobox 1 (Dlx1), mRNA	2.5
136	AF257303	Mus musculus synaptotagmin II (Syt2) gene, complete cds	0.85
137	Z92837	Caenorhabditis elegans cosmid R03E1, complete sequence	0.027
138	AF258528	Loligo pealei phospholipase C mRNA, complete cds	0.74
139	M65252	Bacillus thuringiensis alesti delta endotoxin gene, complete cds	0.075
140	M14954	D.melanogaster (W-IR1 mutation) I factor DNA, complete cds	0.053
141	Z48636	C.novyi gene for alpha-toxin	0.026
142	XM_036491	Homo sapiens hypothetical protein MGC3771 (MGC3771), mRNA	0.002

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
143	AK020198	Mus musculus 15 days embryo male testis cDNA, RIKEN full-length enriched library, clone:8030442B05, full insert sequence	0.53
144	AJ404228	Saccharomyces douglasii mitochondrial tRNA gene cluster	0.022
145	AJ297708	Rattus norvegicus RT6 gene for T cell differentiation marker RT6.2, exons 1-8	0.024
146	AK026697	Homo sapiens cDNA: FLJ23044 fis, clone LNG02454	1.9
147	AE001370	Plasmodium falciparum chromosome 2, section 7 of 73 of the complete sequence	0.017
148	AF325208	Acetabularia acetabulum nicotinamide nucleotide transhydrogenase (Nnt) gene, exons 6, 7, 8, and complete cds	5.2
149	L78770	Homo sapiens (subclone 1_f7 from P1 H43) DNA sequence	0.068
		Glycine max glutamine synthetase gamma 2 subunit gene, 5'-flanking	
150	AF182214	region and partial cds	0.2
		Lolium rigidum clone LR1 acetyl-CoA carboxylase mRNA, partial	
151	AF359513	cds; nuclear gene for plastid product	0.2
		Pseudomonas aeruginosa PA01, section 359 of 529 of the complete	
152	AE004798	genome	2.6
153	NM_021699	Rattus norvegicus serine/threonine kinase (LOC60328), mRNA	1.8
154	XM_040539	Homo sapiens interphotoreceptor matrix proteoglycan 200 (SPACRCAN), mRNA	2.3
155	AE006894	Sulfolobus solfataricus section 253 of 272 of the complete genome	0.027
156	Y14603	Erwinia amylovora srlA, srlE, srlB, srlD, srlM and srlR genes	0.8
157	AF181720	Homo sapiens RU2AS (RU2) gene, complete cds; and RU2S (RU2) gene, partial cds	0.051
158	XM_043815	Homo sapiens hypothetical protein FLJ21736 (FLJ21736), mRNA	1.1
159	AE006059	Pasteurella multocida PM70 section 26 of 204 of the complete genome	1.6
160	AF151097	Homo sapiens voltage-dependent anion channel (VDAC1) gene, exons 8 and 9 and complete cds	0.074
161	AF165136	Mycoplasma mycoides mycoides LC hypothetical surface located membrane protein and lipoprotein B precursor (lppB) genes, partial cds	0.23
162	XM_016392	Homo sapiens hypothetical protein FLJ12768 (FLJ12768), mRNA	2.1
163	AF170544	Magnaporthe grisca vacuolar-ATPase (VATP) mRNA, complete cds	0.15
164	AY045680	Arabidopsis thaliana At1g08350/T27G7_4 mRNA, complete cds	0.009
165	AF285100	Bos taurus lysophosphatidic acid acyltransfesrase gene, complete cds	1.6
166	AF194170	Dictyostelium discoideum cudA protein gene, promotor sequence and partial cds	0.039
167	AF242446	Homo sapiens LINE-1 insertion dimorphism LID-3 empty site sequence	0.043

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Pan troglodytes gamma-aminobutyric acid receptor A5 subunit	
168	AF084196	duplicated gene, 5'UTR region	0.88
169	AK021754	Homo sapiens cDNA FLJ11692 fis, clone HEMBA1004983	0
170	AE006002	Caulobacter crescentus section 328 of 359 of the complete genome	1.6
171	AL096734	Homo sapiens mRNA; cDNA DKFZp434M011 (from clone DKFZp434M011)	0.015
		Homo sapiens 2'-5'oligoadenylate synthetase 3 (OAS3) gene, promoter	
172	AF251351	region	2.7
		Agrocybe aegerita B type DNA polymerase (Mtpol) gene, complete	
		cds; tRNA-Asn gene, complete sequence; and unknown genes,	
173	AF061244	mitochondrial genes for mitochondrial products	0.0006
174	X95275	P.falciparum complete gene map of plastid-like DNA (IR-A)	0.1
176	M88599	Entamoeba histolytica P-glycoprotein-1 (pgp1) gene, complete cds	0.12
		Oryza sativa microsatellite MRG1949 containing (AT)X40, closest to	
177	AY019624	marker R1167, genomic sequence	0.017
178	NM_026040	Mus musculus RIKEN cDNA 2810036K01 gene (2810036K01Rik), mRNA	e-101
179	AL162040	Homo sapiens mRNA; cDNA DKFZp434N199 (from clone DKFZp434N199)	e-152
180	AJ295190	Human coxsackievirus A11 genomic RNA for partial polyprotein gene, isolate VR-169, ATCC	1.5
181	AJ298684	Phylloxera sp. MBLM2 mitochondrial ATP6 gene for ATP synthase A chain subunit 6	0.51
		Candidatus Carsonella ruddii natural-host Calophya schini RNA	
		polymerase beta subunit (rpoB) and RNA polymerase beta-prime	
183	AF268064	subunit (rpoC) genes, partial cds	0.012
184	U27374	Human Menkes disease gene (ATP7A), exon 16	0.51
		Epizootic haemorrhagic disease virus serotype 1 (EHDV-1) gene	
185	D10767	encoding VP2 protein	0.049
186	AJ002256	Mus musculus minisatellite tandem repeat (MMS73)	0.058
187	AB007546	Homo sapiens gene for LECT2, complete cds	2E-11
188	AF254573	Cricetulus griseus origin recognition complex subunit 2 mRNA, complete cds	0.57
		Homo sapiens cDNA: FLJ21250 fis, clone COL01253, highly similar	
1		to AB020527 Homo sapiens mRNA for Na/PO4 cotransporter	
189	AK024903	homolog	e-135
		Lolium rigidum clone LR1 acetyl-CoA carboxylase mRNA, partial	
190	AF359513	cds; nuclear gene for plastid product	0.15
		Mus musculus mRNA for hypothetical protein expressed in	l
191	AJ237585	thymocytes (clone MFT,M05.13/MTA,B10.066), partial	4.4
192	AF233591	Mus musculus son-of-sevenless 1 (Sos1) gene, partial cds	0.52
193	X81190	M.musculus DNA flanking site of transgene insertion	4.8
194	AF232246	Tetrahymena thermophila clone Tlr1 Int integrase-like protein gene,	0.53
174	MFZ3ZZ40	complete cds	0.33

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	
NU	ACCESSN		P VALUE
225	XM_042689	Homo sapiens fasciculation and elongation protein zeta 2 (zygin II) (FEZ2), mRNA	0.69
	4 TO 1 WOOD	Anoplostoma viviparum isolate Brh10 large subunit ribosomal RNA	
226	AF317083	gene, partial sequence; mitochondrial gene for mitochondrial product	0.22
227	177006426	Lactococcus lactis subsp. lactis IL1403 section 198 of 218 of the	
227	AE006436	complete genome	0.029
220	177025505	Homo sapiens cDNA FLJ14629 fis, clone NT2RP2000448, weakly	
228	AK027535	similar to KES1 PROTEIN	7E-76
220	DC004200	Homo sapiens, NIMA (never in mitosis gene a)-related kinase 6, clone	
229	BC004209	MGC:4434 IMAGE:2958695, mRNA, complete cds	3E-36
200	4.0000100	Homo sapiens (subclone 2_h8 from BAC H111) DNA sequence,	
230	AC002183	complete sequence	0.0000004
		Prochlorococcus marinus UreC (ureC), UreB (ureB), UreA (ureA),	
		UreD (ureD), UreE (ureE), UreF (ureF), and UreG (ureG) genes,	
231	AF242489	complete cds	0.0003
232	AF385396	Homo sapiens psoriasis susceptibility gene, partial sequence	0.027
233	AE006749	Sulfolobus solfataricus section 108 of 272 of the complete genome	2.1
234	X73501	H.sapiens gene for cytokeratin 20	0.25
		Glyptotendipes pallens 18S rRNA gene (partial), 2S rRNA gene, 28S	
		rRNA gene (partial), internal transcribed spacer 1 (ITS1) and internal	
235	AJ296801	transcribed spacer 2 (ITS2), clone Gpaits9	0.49
236	X17627	C. thummi DNA transposable element TECth1	0.59
		Eutamias dorsalis cytochrome b (cytb) gene, complete cds;	
237	AF157924	mitochondrial gene for mitochondrial product	0.19
		Homo sapiens similar to calcium channel, voltage-dependent, alpha	
238	XM_040836	1H subunit (H. sapiens) (LOC91809), mRNA	0.38
		Oreina melanocephala 16S large subunit ribosomal RNA gene, partial	
239	AF097103	sequence; mitochondrial gene for mitochondrial product	0.23
		Mus musculus 13 days embryo head cDNA, RIKEN full-length	
240	AK013974	enriched library, clone:3110002D09, full insert sequence	0.73
241 242	AK022073	Homo sapiens cDNA FLJ12011 fis, clone HEMBB1001653	0.26
242	XM_012392 AB048604	Homo sapiens ubiquitin specific protease 8 (USP8), mRNA	2
243	AB048604	Mus musculus Neu-2 mRNA for sialidase, complete cds	0.54
		Drepanaphis utahensis 12S small subunit ribosomal RNA gene,	
		partial sequence; tRNA-Val gene, complete sequence; and 16S large	
		subunit ribosomal RNA gene, partial sequence; mitochondrial genes	
244	AF275235	for mitochondrial products	0.076
245	XM 032956	Homo sapiens cat eye syndrome chromosome region, candidate 2 (CECR2), mRNA	2.2
- 10		Homo sapiens secretory protein SEC8; KIAA1699 protein (SEC8),	
246	XM 050246	mRNA	0
247	X69200	D.tigrina Dth-1 gene, exon 2	0.49
248	M33582	O.cuniculus beta-casein gene, complete cds	0.006
250	AB048604	Mus musculus Neu-2 mRNA for sialidase, complete cds	0.46

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
251	U48594	Heliothis virescens insect storage protein (SP4) mRNA, partial cds	0.022
1		Human DNA sequence from clone RP1-250J21 on chromosome	
252	AL162583	Xp11.22-11.4, complete sequence [Homo sapiens]	0.2
253	AF284038	Cucurbita maxima phloem serpin-1 mRNA, complete cds	0.19
254	AK026699	Homo sapiens cDNA: FLJ23046 fis, clone LNG02491	0.6
		Trioxys betulae NADH dehydrogenase 1 gene, mitochondrial gene	
255	AF069199	encoding mitochondrial protein, partial cds	0.19
256	Z12028	L. pimpinellifolium gene encoding vacuolar invertase	0.7
		Panonychus mori mitochondrial CO1 gene for cytochrome oxidase	
257	AB041256	subunit 1, partial cds, strain: TtDe	0.21
258	AK022820	Homo sapiens cDNA FLJ12758 fis, clone NT2RP2001328	0.25
259	AE006782	Sulfolobus solfataricus section 141 of 272 of the complete genome	1.4
260	AF027656	Homo sapiens cholesteryl ester transfer protein gene, promoter region	0,00004
		H. sapiens CpG island DNA genomic Mse1 fragment, clone 22b3,	
261	Z57957	reverse read cpg22b3.rt1a	0.00003
1 1		Orthetrum cancellatum large subunit ribosomal RNA gene, partial	1
		sequence; tRNA-Valine gene, complete sequence; and small subunit	
		ribosomal RNA gene, partial sequence; mitochondrial genes for	l
262	AF266097	mitochondrial products	0.077
263	XM_003119	Homo sapiens dynein light chain-A (LOC51143), mRNA	5.6
264	AF099990	Rattus norvegicus Ste-20 related kinase SPAK mRNA, complete cds	0.54
204	A1-033330	Rattus not vegicus Sie-20 Telateu kinase SFAR nikeva, complete cus	0.54
265	AE007394	Streptococcus pneumoniae section 77 of 194 of the complete genome	2.2
203	AL007334	Leishmania tarentolae kinetoplast mitochondrial MURF2 edited	2.2
266	L07545	mRNA, complete cds	0.002
200	207343	Homo sapiens complement component 2 (C2) gene allele b, exons 1	0.002
267	L09706	through 8	0.073
207	105700	Pauesia unilachni 16S ribosomal RNA gene, partial sequence;	0.075
268	AF174326	mitochondrial gene for mitochondrial product	0.063
269	Z79794	M.musculus dystrobrevin gene, exon 8	1.8
-207	217171	Oryza sativa gene for water channel protein RWC3, promoter region	1.0
270	AB029325	and complete cds	1.7
	- 12027020	Xenopus laevis neural specific DNA binding protein (Xgli3) mRNA.	
271	U42461	complete cds	1.6
272	XM 029784	Homo sapiens CGI-53 protein (LOC51098), mRNA	6E-23
273	D14011	Mouse reg II gene for regenerating protein II, complete cds	0.071
		Homo sapiens short-chain dehydrogenase/reductase 1 (SDR1) gene,	
274	AF179235	exons 2, 3, and 4	2.1
		Li	

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		Land Control of the C	
NO	ACCESSN	DESCRIP	P VALUE
275	AF264911	Campylobacter jejuni putative UDP-N-acetylmuramate-alanine ligase (murC) gene, partial cds, putative integral membrane protein, putative MutS (mutS), restriction and modification enzyme Cjel (cjel), putative transferase, and putative integral membrane >	
276	AF266074	Gomphus exilis large subunit ribosomal RNA gene, partial sequence, iRNA-Valine gene, complete sequence; and small subunit ribosomal RNA gene, partial sequence; mitochondrial genes for mitochondrial products	0.021
277	AF229988	Cryptosporidium parvum P-type ATPase3 gene, complete cds	1.9
278	AF250346	Xenopus laevis early growth response protein 1 (egr1) gene, partial cds	0.25
279	AL157448	Homo sapiens mRNA; cDNA DKFZp761E0311 (from clone DKFZp761E0311)	0.00001
280	AF386078	Homo sapiens serine-cysteine proteinase inhibitor clade C member 1 (SERPINC1) gene, complete cds	2.1
281	S80932	MVAT5-RX2 VSG=variant surface glycoprotein {promoter} [Trypanosoma brucei=African trypanosomes, Genomic, 6583 nt]	0,003
282	U09478	Dictyostelium discoideum KAx-3 LagC protein (lagC) mRNA,	0.000
282	U63580	complete cds Oxytricha fallax micronuclear actin I gene, partial sequence	0.008
284	AF348411	Macaca mulatta Per4 pseudogene sequence	0.0022
204	Ar546411	Homo sapiens cellular apoptosis susceptibility protein (CSE1) gene.	0.0008
285	AF053645	exons 3 through 10	0.0002
287	U90928	Arabidopsis thaliana głyoxalase II mitochondrial isozyme (Glx2-1) mRNA, nuclear gene encoding mitochondrial protein, complete cds Carabus pseudopusio mitochondrial ND5 gene for NADH	2.2
288	AB050751	dehydrogenase subunit 5, partial cds	1.6
289	AF275220	Prociphilus fraxinifolii 12S small subunit ribosomal RNA gene, partial sequence; tRNA-Val gene, complete sequence; and 16S large subunit ribosomal RNA gene, partial sequence; mitochondrial genes for mitochondrial producis	0.003
290	NM_014358	Homo sapiens C-type (calcium dependent, carbohydrate-recognition domain) lectin, superfamily member 9 (CLECSF9), mRNA	0.009
291	D25274	Homo sapiens mRNA, clone:PO2ST9	1E-38
292	AF111936	Callithrix jacchus isolate MJ50037 alpha (1,2) fucosyltransferase (FUT1) gene, partial cds	1.8
293	AL592159	Human DNA sequence from clone RP13-237P14 on chromosome X, complete sequence [Homo sapiens]	0.18
294	AK013928	Mus musculus 13 days embryo head cDNA, RIKEN full-length enriched library, clone:3100003M19, full insert sequence	1.9
295	AF289199	Mus musculus junctin and aspartyl beta-hydroxylase (Asph) genes, partial cds	0.079

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			l
NO	ACCESSN	DESCRIP	P VALUE
296	AE001425	Plasmodium falciparum chromosome 2, section 62 of 73 of the complete sequence	0,008
297	XM 011029	Homo sapiens LIM domain-containing preferred translocation partner in lipoma (LPP), mRNA	0.23
298	AF105034	Arabidopsis thaliana delta7 sterol C-5 desaturase (STE1) gene, complete cds	0,009
299	L77040	Homo sapiens (subclone 8 cl 1 from P1 H22) DNA sequence	9E-14
		Tome suprems (outcome o_orr resmit r range) 2 thristochasine	72 1.
300	AF278536	Parastrongyloides trichosuri heat shock 70 protein gene, complete cds	0.61
302	Z28216	S.cerevisiae chromosome XI reading frame ORF YKL216w	1.7
303	AJ008049	Chrysolina colasi 16S rRNA gene	0.0007
304	AF280812	Glycine max putative Hs1pro-1-like receptor mRNA, complete cds	0.67
		Mus musculus 10 day old male pancreas cDNA, RIKEN full-length	
305	AK007831	enriched library, clone:1810048N21, full insert sequence	0.047
1		Human DNA sequence from clone RP11-424E21 on chromosome 13,	
306	AL163543	complete sequence [Homo sapiens]	0.64
307	AF045022	Bos taurus phosphatidic acid-preferring phospholipase A1 mRNA, complete cds	5
308	AB052731	Oikopleura longicauda OilBra mRNA for brachyury protein, complete cds	1.8
		Holbrookia maculata clone HMFL 12S ribosomal RNA gene, partial	
309	AF194275	sequence; mitochondrial gene for mitochondrial product	6.3
310	AJ400814	Dictyostelium discoideum srfA gene, alternative promoters	0.024
		Mus musculus adult male kidney cDNA, RIKEN full-length enriched	
311	AK002854	library, clone:0610039P13, full insert sequence	0.68
312	U25177	Helobdella robusta cyclin A mRNA, partial cds	0.023
		Mus musculus 13 days embryo liver cDNA, RIKEN full-length	
313	AK011102	enriched library, clone:2510044F14, full insert sequence	0.51
314	AK002181	Homo sapiens cDNA FLJ11319 fis, clone PLACE1010293	0.0006
315	U67282	Human mis5 homolog (MCM6) gene, exon x	0.014
316	XM_040357	Homo sapiens chromosome 6 open reading frame 5 (C6orf5), mRNA	1.5
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
317	AK006073	library, clone:1700017M07, full insert sequence	6.7
318	AE001381	Plasmodium falciparum chromosome 2, section 18 of 73 of the complete sequence	0.18
319	NM_017678	Homo sapiens hypothetical protein FLJ20127 (FLJ20127), mRNA	3E-57
		Mus musculus, Similar to coxsackicvirus and adenovirus receptor,	
320	BC004680	clone MGC:5878 IMAGE:3500491, mRNA, complete cds	0.24
321	AK002181	Homo sapiens cDNA FLJ11319 fis, clone PLACE1010293	0.007
	******	Dictyostelium discoideum myosin heavy chain kinase B (MHCK-B)	
322	U90946	mRNA, complete cds	0.003
323	Y14277	Drosophila melanogaster mRNA for nuclear protein SA	0.062

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		Table of Treatest freighton (Diabet Vs. Geneaux)	
NO	ACCESSN	DESCRIP	P VALUE
		H. sapiens DNA segment containing (CA) repeat; clone AFM248wc1;	T TILBOD
324	Z17272	single read	0.25
		E.gracilis chloroplast P-700 chlorophyll alpha apoprotein (psaA and	
325	M37526	psaB) genes, complete cds	0.025
326	Y14952	Mus musculus gene encoding immunoglobulin J chain precursor	0.24
		Streptococcus pyogenes M1 GAS strain SF370, section 125 of 167 of	
327	AE006596	the complete genome	6.8
		Candida albicans N-acetyl-glucosamine-6-phosphate deacetylase	-
		(DAC1) and glucosamine-6-phosphate deaminase (NAG1) genes,	
		complete cds; and hexokinase (HXK1) pseudogene, complete	
328	AF079804	sequence	2
		Homo sapiens Machado-Joseph disease (spinocerebellar ataxia 3,	
200	7D 5 000015	olivopontocerebellar ataxia 3, autosomal dominant, ataxin 3) (MJD), mRNA	
329	XM_029315 AK022820	Homo sapiens cDNA FLJ12758 fis, clone NT2RP2001328	0.24
330	AN.022020	Mus musculus heat shock transcription factor 1 (Hsf1) gene, partial	0.24
331	AF059275	cds	0.085
331	PHOSSETS	Homo sapiens mRNA; cDNA DKFZp564F133 (from clone	0.003
332	AL049263	DKFZp564F133)	0.059
-552	122019200	212 2500 12 125)	0.003
333	AP001233	Homo sapiens genomic DNA, chromosome 2p11.2, clone:cos607/2	0.003
334	AC017021	Homo sapiens BAC clone RP11-208M4 from 7, complete sequence	0.0000005
335	U03248	Human chromosome 21 clone pVC1.23c	3E-56
336	XM_046758	Homo sapiens tensin (TNS), mRNA	2
337	AJ279150	Homo sapiens partial TFNR gene for transcription factor-like nuclear regulator, exon 31	0
338	AF282896	Rattus norvegicus chemokine RANTES gene, promoter sequence	0.001
339	AK000939	Homo sapiens cDNA FLJ10077 fis, clone HEMBA1001864	0.21
	17100517	20.11	0.000
340	AF188517 AF202562	Staphylococcus aureus Blt-like protein SbtA (sbtA) gene, complete cds	0.008
341	AF202362	Homo sapiens DNA methyltransferase (DNMT1) gene, exon 13	0.22
342	XM 017967	Homo sapiens hypothetical protein FLJ13732 similar to tensin (FLJ13732), mRNA	2.7
343	AF051685	Homo sapiens clone SBS-12 SATB1 binding sequence in vivo	0.21
343	711 05 1005	Campylobacter icjuni argininosuccinate lyase (argH) gene, complete	0.22
344	M77188	cds	0.22
345	U95137	Rattus norvegicus Na+-Ca+ exchanger (NCX1) gene, exon 1-Ht	0.24
		Mus musculus 6 days neonate head cDNA, RIKEN full-length	
346	AK019943	enriched library, clone:5430403G16, full insert sequence	0.74
		Mus musculus, ubiquitin specific protease 14, clone MGC:7106	
347	BC005571	IMAGE:3157723, mRNA, complete cds	0.41
			100.46
348	XM_026968	Homo sapiens hypothetical protein MGC2668 (MGC2668), mRNA	4E-16

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Table 3A Nearest Neighbor (BlastN vs. Genbank) SEQ ID NO ACCESSN DESCRIP P VALUE Homo sapiens cDNA: FLJ21250 fis, clone COL01253, highly similar to AB020527 Homo sapiens mRNA for Na/PO4 cotransporter 349 AK024903 7E-28 Oryctolagus cuniculus fructose 1,6, bisphosphate aldolase (AldB) 350 U85645 gene, complete cds 0.075

000	0.00 0.10	gene, comprete cus	0.012
		Homo sapiens fibrinogen, B beta polypeptide (FGB) gene, complete	
351	AF388026	cds	5.8
		Homo sapiens pro-melanin-concentrating hormone-like 2 protein	
352	AY028321	(PMCHL2) gene, exons 4, 5a, and 5b, alternatively spliced	0.0003
		Human DNA sequence from clone RP11-79A21 on chromosome X,	
353	AL513491	complete sequence [Homo sapiens]	1.9
		Chimpanzee retrovirus-like sequence-isoleucine b (RTVL-Ib) gene, 5'	
354	M92068	and 3' LTR	1.5
355	XM_002120	Homo sapiens zinc finger protein 262 (ZNF262), mRNA	0.007
356	X02435	Tetrahymena mitochondrial gene for tRNA-Phe (GAA)	0.028
357	XM_033459	Homo sapiens syntaxin binding protein 1 (STXBP1), mRNA	2.1
358	NM_015770	Mus musculus non-agouti (a), mRNA	0.64
		Staphylococcus aureus repNVH99 gene for replication protein and	
359	AJ296103	smr gene	0.15
360	Z12840	O.cuniculus mRNA for protein of unknown function	e-131
		Homo sapiens hypothetical gene supported by AK001938; AK001941	
361	XM 038370	(LOC91424), mRNA	0.0002
		Rhipidomys mastacalis cytochrome B (cytB) gene, mitochondrial gene	
362	AF108684	encoding mitochondrial protein, partial cds	0.009
363	NM 020045	Mus musculus HIRA-interacting protein 5 (HIRIP5), mRNA	1.8
364	AK024439	Homo sapiens mRNA for FLJ00029 protein, partial cds	0.73
365	AK025445	Homo sapiens cDNA: FLJ21792 fis, clone HEP00441	2.1
	122020110	Homo sapiens genomic DNA, chromosome 21g22.1, D21S226-AML	
366	AP000307	region, clone:f6C7, complete sequence	0.00004
500	1200000	Methanobacterium thermoautotrophicum from bases 264585 to	0,000
367	AE000818	276866 (section 24 of 148) of the complete genome	2.3
368	AK001164	Homo sapiens cDNA FLJ10302 fis, clone NT2RM2000042	0.13
369	XM 029101	Homo sapiens KIAA0947 protein (KIAA0947), mRNA	0.95
370	AK022191	Homo sapiens cDNA FLJ12129 fis, clone MAMMA1000198	0.029
371	Z54280	S. scrofa gene for skeletal muscle ryanodine receptor	1.9
3/1			
372	AY037259	Arabidopsis thaliana AT3g52120/F4F15 230 mRNA, complete cds	9.6
372		Arabidopsis thaliana AT3g52120/F4F15_230 mRNA, complete cds Homo sapiens acid sphingomyelinase-like phosphodiesterase	9.6
	AY037259	Homo sapiens acid sphingomyelinase-like phosphodiesterase	9.6
372 373			
	AY037259	Homo sapiens acid sphingomyelinase-like phosphodiesterase (ASM3A), mRNA	
373	AY037259 XM_034599	Homo sapiens acid sphingomyelinase-like phosphodiesterase (ASM3A), mRNA Homo sapiens SWI/SNF related, matrix associated, actin dependent	0
	AY037259	Homo sapiens acid sphingomyelinase-like phosphodiesterase (ASM3A), mRNA Homo sapiens SWJ/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily f, member 1 (SMARCFI), mRNA	
373 374	AY037259 XM_034599 XM_046914	Homo sapiens acid sphingomyelinase-like phosphodiesterase (ASM3A), mRNA Homo sapiens SWJSNF related, matrix associated, actin dependent regulator of chromatin, subfamily f, member 1 (SMARCF1), mRNA Mycoplasma-like sp. 235 and 16S ribosomal RNAs, Tyr-transfer RNA	0 3E-16
373	AY037259 XM_034599	Homo sapiens acid sphingomyelinase-like phosphodiesterase (ASM3A), mRNA Homo sapiens SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily f, member 1 (SMARCFI), mRNA Mycoplasma-like sp. 23S and 16S ribosomal RNAs, Tyr-transfer RNA and Ile-transfer RNA, complete cds	0
373 374	AY037259 XM_034599 XM_046914	Homo sapiens acid sphingomyelinase-like phosphodiesterase (ASM3A), mRNA Homo sapiens SWJSNF related, matrix associated, actin dependent regulator of chromatin, subfamily f, member 1 (SMARCF1), mRNA Mycoplasma-like sp. 235 and 16S ribosomal RNAs, Tyr-transfer RNA	0 3E-16

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
377	AF106915	Phodopus sungorus luteinizing hormone beta subunit mRNA, partial cds	0.61
		Human DNA sequence from clone RP1-316I5 on chromosome 20	
		Contains the 3' end of the KIF3B gene encoding a kinesin family	
378	AT 251900	member 3B (KIAA0359) and ESTs, complete sequence [Homo sapiens]	
3/8	AL354800	Homo sapiens mRNA full length insert cDNA clone EUROIMAGE	0.0001
379	AL133430	233360	0,66
380	AF085894	Homo sapiens full length insert cDNA clone YP97D11	0.0002
		Sesbania rostrata mRNA for phosphoenolpyruvate carboxylase (pepc	
381	AJ286750	gene)	0.068
		Homo sapiens region containing hypothetical protein; mutL (E. coli)	
382	XM_040415	homolog 3 (LOC82389), mRNA	e-128
383	Y09794	A.rabiei microsatellite DNA	1.2
384	AB047905	Macaca fascicularis brain cDNA, clone:QnpA-15911	1.8
		Homo sapiens cDNA FLJ10589 fis, clone NT2RP2004389, weakly	
	*****	similar to PROBABLE MITOCHONDRIAL 40S RIBOSOMAL	1 1
385	AK001451	PROTEIN S9 PRECURSOR	1.5
386	AF010473	Schizosaccharomyces pombe myosin-like protein Sp8 (sp8) mRNA, partial cds	0.19
380	A1010473	Homo sapiens mRNA; cDNA DKFZp434G1615 (from clone	0.19
388	AL133604	DKFZp434G1615)	0,00006
		Homo sapiens methyl-CpG binding domain protein 3 (MBD3),	
389	XM_009252		0.055
		Polyandrocarpa misakiensis mRNA for putative eukaryotic petide	
390	AB053116	chain release factor subunit 1, complete cds	1.5
391	AL138666	S.pombe chromosome I cosmid c694	1,3
		Pleurotus ostreatus small subunit ribosomal RNA gene, partial	
392	AF091933	sequence; mitochondrial gene for mitochondrial product	0.078
393	Z37964	Staphylococcus sp. genes encoding QacC and replication protein (rep827)	0.073
393	AB020712	Homo sapiens mRNA for KIAA0905 protein, complete cds	2E-64
354	AB020712	Mus musculus, Similar to prostaglandin E receptor 4 (subtype EP4),	26-04
395	BC009023	clone MGC:7183 IMAGE:3481696, mRNA, complete cds	1.5
	2000	Homo sapiens odz (odd Oz/ten-m, Drosophila) homolog 1 (ODZ1),	7.0
396	XM_010128	mRNA	0.55
397	XM 011386	Homo sapiens hypothetical protein PRO0813 (PRO0813), mRNA	5.3
397	XM 040694	Homo sapiens AD031 protein (AD031), mRNA	1.3
320	25LVI_0-10094	Pasteurella multocida PM70 section 144 of 204 of the complete	1.3
399	AE006177	genome	5
		Botrytis cinerea strain T4 cDNA library under conditions of nitrogen	
400	AL113079	deprivation	1.2
401	U39694	Mycoplasma genitalium section 16 of 51 of the complete genome	0.84
402	XM_016314	Homo sapiens ankyrin 2, neuronal (ANK2), mRNA	4.6

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Homo sapiens likely homolog of rat kinase D-interacting substance of	
403	XM_045362	220 kDa; KIAA1250 protein (KIAA1250), mRNA	7E-77
404	NM_028513	Mus musculus RIKEN cDNA 1700052K15 gene (1700052K15Rik), mRNA	9
		Solanum allophyllum NADH dehydrogenase subunit (ndhF) gene,	
405	U47416	chloroplast gene encoding chloroplast protein, partial cds	0.2
406	NC 002502	Lactococcus lactis subsp. lactis plasmid pCi305, complete sequence	5.3
-400	110_002302	Staphylococcus epidermidis strain SR1 clone step.1010e06 genomic	3.5
407	AF269561	sequence	0.19
409	AF336128	Lumpy skin disease virus strain Neethling, 5' partial sequence	1.5
410	AB058397	Oryza sativa CHS gene for chalcone synthase, complete cds	0.16
710	AD030377	oryza sauva crio gene ioi cinacone synniase, comprete cus	0.10
411	AE006830	Sulfolobus solfataricus section 189 of 272 of the complete genome	0.21
711	ALGOGOGO	Arabidopsis thaliana unknown protein (T20O10 100/AT3g63000)	0.21
412	AY035064	mRNA, complete cds	0.22
712	A1033004	Platybrachys decemmacula 16S ribosomal RNA gene, partial	0.22
413	AF158049	sequence; mitochondrial gene for mitochondrial product	0.063
414	NM 026444	Mus musculus citrate synthase (Cs), mRNA	1.7
415	BC007820	Homo sapiens, clone IMAGE:4303165, mRNA	0
413	DC007620	Caenorhabditis elegans mRNA for MAGUK protein DLG-1 (dlg-1	
416	AJ295228	gene)	0.2
		Flammulina velutipes mitochondrial gene for DNA polymerase, RNA	
417	AB028633	polymerase, complete and partial cds	0.6
418	Y09794	A.rabiei microsatellite DNA	1.2
419	AB054063	Pagrus major lpl gene for lipoprotein lipase, complete cds	5.9
420	AF323928	Plasmodium falciparum GcpE (gcpE) gene, complete cds	0.14
421	M31012	Chicken MHC class I B-FTV-B12 alpha-chain gene, complete cds	1.2
422	Z49806	B.taurus mRNA for adenylyl cyclase type VII	6.1
		Simulation and a ration and a system to the ration and a ration and a system to the ration and a syste	
423	AB009907	Luciola kuroiwae mitochondrial DNA for 16S rRNA, partial sequence	0.003
424	X76693	C.coli (UA585) sodB gene for superoxide dismutase	0.45
		Carex capitata tRNA-Thr, partial sequence; trnT-trnL intergenic	
		spacer, tRNA-Leu, and trnL-trnF intergenic spacer, complete	
]		sequence; and tRNA-Phe, partial sequence; chloroplast genes for	
425	AF284942	chloroplast products	0.29
		Mus musculus adult male thymus cDNA, RIKEN full-length enriched	
426	AK017986	library, clone;5830443G21, full insert sequence	5,1
427	AL512549	S.pombe chromosome I BAC pB2B4	1.6
428	AF283669	Homo sapiens kallikrein 14 (KLK14) gene, complete cds	2.3
429	Z12842	O.cuniculus mRNA for protein of unknown function	5E-73
430	XM_016894	Homo sapiens LOC86241 (LOC86241), mRNA	3E-65
431	X96616	P.primaurelia gene encoding 156D surface antigen	0.23
432	X02175	Schizosaccharomyces pombe cdc10 start gene	0.027
433	AK021807	Homo sapiens cDNA FLJ11745 fis, clone HEMBA1005526	0.001

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		I .	
NO	ACCESSN	DESCRIP	P VALUE
		Homo sapiens protein phosphatase 2 (formerly 2A), regulatory subunit	
434	XM_005233	B (PR 52), alpha isoform (PPP2R2A), mRNA	5E-09
435	X52623	Rice 4-CL gene for 4-coumarate-CoA ligase (EC 6.2.1.12)	0.63
		Mus musculus adult male cerebellum cDNA, RIKEN full-length	
436	AK005294	enriched library, clone:1500019O16, full insert sequence	1E-16
437	AE002337	Chlamydia muridarum, section 65 of 85 of the complete genome	0.049
438	AF106574	Caenorhabditis elegans cosmid E02D9, complete sequence	0.7
439	X56866	S.alba 5S rRNA gene	0.74
		Drosophila mauritiana Cu-Zn superoxide dismutase (Sod) gene, exons	
440	AF127158	1 and 2 and complete cds	1.7
441	U08466	Nephila clavipes alanine tRNA gene, complete sequence	2.1
		Oryza sativa microsatellite MRG3561 containing (TA)X22, genomic	
442	AY021236	sequence	1.5
443	L09262	Arabidopsis thaliana phytochrome b gene, exons 2 and 3	0.079
444	AC007095	Homo sapiens BAC clone RP11-303A22 from 7, complete sequence	0.23
		Mus musculus 11 days embryo cDNA, RIKEN full-length enriched	
445	AK012607	library, clone:2700094F01, full insert sequence	0.19
446	AF309415	Cyprinus carpio ovarian fibroin-like substance-2 mRNA, partial cds	0.008
447	X03956	Potato (tetraploid variety Maris piper) patatin gene	0.003
448	BC008617	Mus musculus, clone IMAGE:3588380, mRNA, partial cds	0.07
		Mauritiella armata 18S rRNa gene (partial), 5.8S rRNA gene, 26S	
		rRNA gene (partial) and internal transcribed spacers 1 and 2 (ITS1,	
449	AJ242146	ITS2), clone 1	0.07
450	M67489	Bovine desmocollin mRNA, complete cds	0.066
451	AB048954	Macaca fascicularis brain cDNA, clone:QnpA-10509	0.48
452	AE001136	Borrelia burgdorferi (section 22 of 70) of the complete genome	0.18
	•	Plasmodium falciparum chromosome 2, section 35 of 73 of the	
453	AE001398	complete sequence	0.025
454	AF218904	Homo sapiens attractin precursor (ATRN) gene, exon 19	1.6
455	AK025078	Homo sapiens cDNA: FLJ21425 fis, clone COL04162	1E-71
456	M11449	Yeast mitochondrial ori2-ori7 region DNA with putative peptide	0.34
		Homo sapiens hypothetical gene supported by AB007970	
457	XM_031156	(LOC90360), mRNA	0.006
1		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
458	AK015180	library, clone:4930422122, full insert sequence	0.007
459	AC090422	Homo sapiens clone RP11-691A12, complete sequence	0.013
460	U06154	Human clone 2004V-I-1 from chromosome 1q terminal region	1E-19
461	AE001123	Borrelia burgdorferi (section 9 of 70) of the complete genome	0.83
		Enterococcus faecalis gph, ydjH, ydjG, ydjI, pbp4 and ydiC, ORF2	
462	Y17797	and ORF3 genes, partial	0.22
		Homo sapiens SDHD gene for small subunit of cytochrome b of	
463	AB026906	succinate dehydrogenase, complete cds	0.042
464	L13942	Human (clone Cos35) glycerol kinase (GK) gene, exons 1-3	0.042
465	U81510	Spodoptera frugiperda caspase-1 mRNA, complete cds	0.041

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
493	X84060	H sapiens TCF11 gene, exon 3-6	0.21
494	AP000418	Arabidopsis thaliana genomic DNA, chromosome 3, P1 clone:MPK17	0.003
495	AJ277428	Scenedesmus obliquus mitochondrial tRNA-Ser, tRNA-Asp, 16S rRNA, 23S rRNA genes, strain KS3-2	0.65
496	L11447	Mycoplasma hyorhinis repeat regions in potential metal binding protein gene region	0.046
497	AY029764	Mus musculus TRAM1 mRNA, complete cds	0.23
498	XM_031173	Homo sapiens parathyroid hormone (PTH), mRNA	0.22
499	AF270168	Staphylococcus epidermidis strain SR1 clone step.1051f02 genomic sequence	0.24
500	AF336131	Lumpy skin disease virus strain Neethling, 3' partial sequence	3.5
501	AF195272	Mus musculus serine/threonine kinase AIE1 gene, complete cds	6.7
502	NC_002322	Laqueus rubellus mitochondrion, complete genome	0.26
503	AK007179	Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:1700112J16, full insert sequence	0.24
504	U32706	Haemophilus influenzae Rd section 21 of 163 of the complete genome	0.68
505	AF247555	Drosophila mauritiana heat shock protein 68 gene, partial cds	1.9
506	NM_011146	Mus musculus peroxisome proliferator activated receptor gamma (Pparg), mRNA	0.024
507	M26940	Mouse beta-casein gene, complete cds	0.002
508	AB003043	Phycomyces blakesleeanus classII chitin synthase (PbCHS1) gene, complete eds	
509	U89348	Human papillomavirus type 16 variant, complete sequence	0.68
511	AF270160	Staphylococcus epidermidis strain SR1 clone step.1051e03 genomic sequence	0.02
512	AF119554	Plasmodium falciparum para-aminobenzoic acid synthetase gene, complete cds	0.078
513	AY014991	Waterstoniella sp. CAM-2000 cytochrome oxidase subunit I (COI) gene, partial cds; mitochondrial gene for mitochondrial product	0.68
514	NM 013037	Rattus norvegicus Fos-responsive gene 1 (St2), mRNA	0.085
515	AF169003	Hepatitis C virus isolate G2aK1 polyprotein gene, complete cds	0.028
516	U23442	Tetrahymena thermophila RR internal deletion sequence	0.25
517	AF381638	Homo sapiens SNP EC11 psoriasis susceptibility gene candidate interval, partial sequence	0,0003
518	NM_023755	Mus musculus Tcfcp2-related transcriptional repressor 1 (Crtr1- pending), mRNA	0.009
519	NM_004706	Homo sapiens Rho guanine nucleotide exchange factor (GEF) 1 (ARHGEF1), mRNA	3.4
520	X67164	P.hybrida T-DNA integration region	0.063
521	AB042275	Canine herpesvirus CICPO gene for infected cell protein 0, complete cds	0.66
522	AB019942	Arabidopsis thaliana gene for sigma factor SigA, complete cds	1.4
722		1	2.7

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID	Loopear		
NO	ACCESSN	DESCRIP	P VALUE
523	AF270382	Staphylococcus epidermidis strain SR1 clone step.4041f02 genomic sequence	0.23
		Carex supina tRNA-Thr, partial sequence; trnT-trnL intergenic	
1 1		spacer, tRNA-Leu, and trnL-trnF intergenic spacer, complete	
		sequence; and tRNA-Phe, partial sequence; chloroplast genes for	
524	AF284897	chloroplast products	0.23
525	AK000493	Homo sapiens cDNA FLJ20486 fts, clone KAT08039	0.17
526	U67535	Methanococcus jannaschii section 77 of 150 of the complete genome	0.6
		Drosophila melanogaster genomic scaffold 142000013385431,	
527	AE002754	complete sequence	1.8
		Caenorhabditis elegans SR protein specfic kinase SPK-1 mRNA,	
528	AF241656	complete cds	1.7
		Caenorhabditis elegans putative potassium channel Slo-2 mRNA.	
529	AF173828	complete cds	0.21
		Escherichia coli O157:H7 EDL933 genome, contig 3 of 3, section 1 of	
530	AE005382	290	2.3
531	Z36154	S.cerevisiae chromosome II reading frame ORF YBR285w	0.68
532	AF054126	Rattus norvegicus polymorphic marker D6UIA6 sequence	0.061
		Mus musculus adult male lung cDNA, RIKEN full-length enriched	
533	AK004780	library, clone:1200015C19, full insert sequence	0.7
		Staphylococcus aureus subsp. aureus strain ATCC31889 leukocidin	
534	AF352555	LukS component gene, partial cds	0.2
		Staphylococcus aureus subsp. aureus strain ATCC31889 leukocidin	
535	AF352555	LukS component gene, partial cds	0.1
536	J05246	Mouse alpha-fetoprotein (AFP) gene, 5' flank	0.024
		Rattus norvegicus mytonic dystrophy kinase-related Cdc42-binding	
537	AF021935	kinase (MRCK) mRNA, complete cds	e-141 0.5
538	Y12573	D.melanogaster Jun and 14-3-3 zeta gene	0.5
		Moraxella catarrhalis UDP-glucose dehydrogenase gene, partial cds; and UDP-glucose 4-epimerase (galE) and L-glutamine:D-fructose-6-	
539	AF248584	phosphate aminotransferase (glmS) genes, complete cds	0.026
339	AL 240304	Homo sapiens mRNA; cDNA DKFZp761D0712 (from clone	0.020
540	AL136554	DKFZp761D0712)	0.089
5.0	TIETOUDO!	Platybrachys decemmacula 16S ribosomal RNA gene, partial	0.007
541	AF158049	sequence; mitochondrial gene for mitochondrial product	0.082
		Oryza sativa microsatellite MRG5943 containing (TTA)X55, closest	
542	AY023618	to marker RG908, genomic sequence	0.00009
		Human DNA sequence from clone RP11-113L12 on chromosome 13,	
543	AL591498	complete sequence [Homo sapiens]	3E-14
544	AF183908	Danio rerio P450 aromatase (cyp19b) mRNA, complete cds	0.069
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
545	AK016658	library, clone:4933405A14, full insert sequence	0.53
		Homo sapiens genomic DNA, chromosome 6p21.3, HLA class I	
546	AB014084	region, Cosmid clone:TY7A5, complete sequence	0.011

WO 02/14500 PCT/US01/25840

		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
547	AF285076	Scleria reticularis tRNA-Leu, partial sequence; trnL-trnF intergenic spacer, complete sequence; and tRNA-Phe, partial sequence; chloroplast genes for chloroplast products	0.19
548	XM_047604	Homo sapiens coproporphyrinogen oxidase (coproporphyria, harderoporphyria) (CPO), mRNA	0.17
549	Z81538	Caenorhabditis elegans cosmid F45H10, complete sequence	0.24
550	AF026800	Fundulus heteroclitus cytochrome P4501A (CYP1A) mRNA, complete cds	1.9
551	XM_051897	Homo sapiens phospholipase A2, group IVA (cytosolic, calcium- dependent) (PLA2G4A), mRNA	9E-08
552	XM_015200	Homo sapiens hypothetical protein FLJ22724 (FLJ22724), mRNA	2.1
553	AF266244	Gillichthys mirabilis adenylate cyclase type v-like protein mRNA, complete cds	6
554	Z21818	H. sapiens carcinoembryonic antigen gene	0.23
555	AE002100	Ureaplasma urealyticum section 1 of 59 of the complete genome	0.023
556	XM_045179	Homo sapiens hypothetical gene supported by AL137724 (LOC92454), mRNA	0.19
557	X16509	Rice alpha-amylase gene	1.8
558	M33753	D.melanogaster crumbs protein mRNA, complete cds	5.4
559	D32022	Human mRNA for T cell receptor V beta 8 CDR3, parcial sequence	1.5
560	AF188893	Homo sapiens guanidinoacetate N-methyltransferase (GAMT) gene, complete cds	0.18
561	AF367310	Arabidopsis thaliana At2g36880/T1J8.6 gene, complete cds	0.11
562	AF207699	Elacis guineensis agamous-like MADS box protein OPMADS1 mRNA, complete cds	0.45
564	AB014586	Homo sapiens mRNA for KIAA0686 protein, partial cds	0.17
565	L77040	Homo sapiens (subclone 8_c11 from P1 H22) DNA sequence	8E-14
566	NM_024942	Homo sapiens hypothetical protein FLJ13490 (FLJ13490), mRNA	0.013
567	U88158	Tetrahymena thermophila micronuclear developmentally eliminated sequence region	0.18
568	AF210744	Nannochloropsis oculata large subunit ribosomal RNA gene, partial sequence	0.057
569	AL590384	Human DNA sequence from clone RP11-349A16 on chromosome Xq22.3-24, complete sequence [Homo sapiens]	2E-15
570	AY034379	Capsicum annuum branched-chain amino acid aminotransferase mRNA, complete cds	1.2
		Campylobacter jejuni cytolethal distending toxin A (cdtA), cytolethal distending toxin B (cdtB), and cytolethal distending toxin C (cdtC)	
571	AF038283	genes, complete cds	0.4
572	X98807	A.thaliana mRNA for peroxidase ATP21a	1.8
573	AF111941	Dictyostelium discoideum AX4 development protein DG1148 (DG1148) gene, complete cds	0.0009

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
574	AE002247	Chlamydophila pneumoniae AR39, section 74 of 94 of the complete genome	1.7
575	AF247555	Drosophila mauritiana heat shock protein 68 gene, partial cds	1.4
576	AJ237599	Gallus gallus mRNA for iroquois homologue-2	0.056
577	NM_030995	Rattus norvegicus Microtubule-associated protein 1a (Map1a), mRNA	0.21
578	XM_027638	Homo sapiens KIAA0240 protein (KIAA0240), mRNA	1.5
579	AF310886	Dictyostelium discoideum RacA (racA) gene, complete cds	0.065
580	AB001090	Homo sapiens gene for H-cadherin, exon 1	0.2
581	BC009101	Mus musculus, Similar to hypothetical protein DKFZp761C121, clone MGC:12146 IMAGE:3710846, mRNA, complete cds	0.0008
582	XM 011473	Homo sapiens serine (or cysteine) proteinase inhibitor, clade B (ovalbumin), member 9 (SERPINB9), mRNA	1.8
302	2101_011475	Galium hypocarpium rps16 gene, chloroplast gene, partial intron	1.0
583	AF004051	sequence	0.068
363	FII 004031	scquence	0.006
584	AE007440	Streptococcus pneumoniae section 123 of 194 of the complete genome	0.066
505	1.05647	Homo sapiens fibroblast growth factor receptor gene (located in the	0.000
585	L25647 AF191079	central MHC) signal peptide and consecutive exon Stealth virus 1 clone 3B632, genomic sequence	0.069
586	AF1910/9		0.009
587	L20465	Onchocerca volvulus nicotinic acetylcholine receptor mRNA, partial cds	0.72
588	NM_008713	Mus musculus nitric oxide synthase 3, endothelial cell (Nos3), mRNA	2_
1		Homo sapiens BCL2/adenovirus E1B 19kD-interacting protein 3	
589	NM_004052	(BNIP3), nuclear gene encoding mitochondrial protein, mRNA	4E-26
		Plasmodium falciparum ADP-ribosylation factor 1 (pARF1) mRNA,	
590	U40228	complete cds	0.009
i I		Homo sapiens mRNA; cDNA DKFZp586B0220 (from clone	
591	AL049435	DKFZp586B0220)	0.008
592	AC000981	Homo sapiens (subclone 2_c6 from P1 H31) DNA sequence, complete sequence	0.68
		Neospora caninum small subunit ribosomal RNA and large subunit	
		ribosomal RNA genes, partial sequences and intergenic region, plastid	
593	AF304319	genes for plastid products	0.009
594	BC006517	Homo sapiens, Similar to PPAR binding protein, clone IMAGE:3546031, mRNA	7E-43
595	AJ249162	Homo sapiens promotor enhancer from ISG20 gene	2.1
596	AJ297708	Rattus norvegicus RT6 gene for T cell differentiation marker RT6.2, exons 1-8	0.024
597	Y11779	R.prowazekii ygjT-like gene and 4 open reading frames	0.078
598	AE005660	Escherichia coli O157:H7 EDL933 genome, contig 3 of 3, section 279 of 290	0.53
1	12005000	Homo sapiens RAB11B, member RAS oncogene family (RAB11B),	0.55
599	XM 041617	mRNA	4.8
600		Homo sapiens cDNA FLJ13103 fis, clone NT2RP3002304	0.25

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIPTION	
NO	ACCESSIN	DESCRIP	P VALUE
1		The Day and the County of the	ł
601	Z69388	Human DNA sequence from cosmid L60G9A, Huntington's Disease Region, chromosome 4p16.3 contains ESTs and a CpG island	0,084
602	U84760	Chromatium vinosum dsr locus, complete sequence	0.084
603	AF122050	Hordeum vulgare limit dextrinase gene, complete cds	1.8
- 003	AI 122030	Mus musculus partial gene for acetyl-CoA carboxylase-alpha,	1.0
604	AJ308471	promoter I	0.022
	113300471	Cixius inflatus 16S ribosomal RNA gene, partial sequence;	0.022
605	AF158032	mitochondrial gene for mitochondrial product	0.008
607	XM 012733	Homo sapiens KIAA1012 protein (KIAA1012), mRNA	0.005
		Human mRNA for protein-tyrosine phosphatase HPTPeta, complete	0.000
608	D37781	cds	0.059
		Homo sapiens (subclone 1 g7 from BAC H76) DNA sequence,	
609	AC002252	complete sequence	7E-11
		Peanut witches'-broom phytoplasma RNA polymerase sigma factor	
610	AF160964	(rpoD) gene, complete cds	0.63
611	U67528	Methanococcus jannaschii section 70 of 150 of the complete genome	0.67
		Justicia caudata tRNA-Leu (trnL) gene, chloroplast gene for	
612	AF063134	chloroplast RNA, partial sequence	1.2
613	U67527	Methanococcus jannaschii section 69 of 150 of the complete genome	1.5
614	AK022455	Homo sapiens cDNA FLJ12393 fis, clone MAMMA1002711	0.000001
[]		Poiretia punctata trnK gene, intron, 3' partial sequence; and maturase	l
615	AF272082	(matK) gene, partial cds; chloroplast genes for chloroplast products	0.075
010	AE110461	Homo sapiens G protein-coupled receptor 57 (GPR57) gene, complete	0.005
616	AF112461	cds Dictyostelium discoideum Rac1B (rac1B) gene, complete cds; racK	0.025
617	AF310884	pseudogene, complete sequence; and unknown genes	5.6
017	AF310004	Plasmodium malariae caseinolytic protease C (clpC) gene, partial cds;	3.0
618	AF348342	apicoplast gene for apicoplast product	0.22
619	AB043965	Danio rerio gene for dharma, promoter and partial cds	0.054
620	NC 001839	Petunia vein clearing virus, complete genome	0.61
621	AK023959	Homo sapiens cDNA FLJ13897 fis, clone THYRO1001706	0.67
622	AF028835	Drosophila pallidosa vermilion (v) gene, complete eds	0.13
		- Total paragon (V) Band, Tanpan (L)	
623	U67598	Methanococcus jannaschii section 140 of 150 of the complete genome	0.072
		Pineapple mealybug wilt associated virus-2 polyprotein (ORF1a) and	
		RNA-dependent RNA polymerase genes, partial eds; and hydrophobic	
]		protein p5, heat shock protein 70, p46, coat protein, diverged coat	
624	AF283103	protein, p20, p22, and p6 genes, complete cds	1.5
625	XM_008161	Homo sapiens hypothetical protein FLJ20345 (FLJ20345), mRNA	1E-15
		Homo sapiens adaptor-related protein complex 4, epsilon 1 subunit	
626	XM 035599	(AP4E1), mRNA	6E-22

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
627	AF143066	Lepidopilum scabrisetum ribosomal protein system 4 (rps4) gene, partial cds	0.61
628	U89283	Biomphalaria glabrata myoglobin gene, complete cds	0.46
- 020	007203	Xenopus laevis zona pellucida A glycoprotein homolog (xIZPA)	0.40
629	U44949	mRNA, complete cds	1.6
630	AE007395	Streptococcus pneumoniae section 78 of 194 of the complete genome	0.15
631	BC006963	Mus musculus, clone IMAGE:3708410, mRNA, partial cds	1.6
632	XM_031120	Homo sapiens peptidylglycine alpha-amidating monooxygenase (PAM), mRNA	4E-40
633	AF316122	Kinosternon arizonense isolate KariAZ2 mitochondrial control region, complete sequence	0.69
634	U67561	Methanococcus jannaschii section 103 of 150 of the complete genome	0.22
635	D88010	Human DNA for ribosomal protein S13, complete cds, U14 small nucleolar RNA, complete sequence	0.24
636	AE006321	Lactococcus lactis subsp. lactis IL1403 section 83 of 218 of the complete genome	2.2
637	AF045595	Carassius auratus clone gf-40 glutamic acid decarboxylase isoform 67 (GAD67) mRNA, complete cds	0.74
638	D38378	Human DNA, novel tandem repeat sequence	0.21
639	AF260819	Plasmodium falciparum CG1 protein gene, complete cds	0.003
640	NC_000857	Ceratitis capitata complete mitochondrial genome	0.23
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
641	AK015383	library, clone:4930444M15, full insert sequence	2.1
642	AF286367	Homo sapiens HMGIY gene, promoter	1.6
643	AK019143	Mus musculus 10 days embryo cDNA, RIKEN full-length enriched library, clone:2600001P13, full insert sequence	0.69
		Plasmodium falciparum chromosome 2, section 58 of 73 of the	
644	AE001421	complete sequence	2
645	D16904	Human HepG2 3' region cDNA, clone hmd3a02	0.21
646	XM_003799	Homo sapiens protocadherin beta 10 (PCDHB10), mRNA	0.69
647	NC_001438	Bean golden mosaic virus A component DNA, complete sequence	0.66
		Mus musculus 13 days embryo head cDNA, RIKEN full-length	
648	AK013971	enriched library, clone:3110001P07, full insert sequence	2
649	AF078880	Prosimulium formosum dopa decarboxylase (DDC) gene, partial cds	0.076
650	AJ009600	Begonia taipeiensis rbcL promoter, clone 16320	0.22
651	AF042834	Homo sapiens phosphodiesterase delta subunit gene, exons 2, 3 and 4	5.8
652	AF244214	Dysdera ratonensis 16S ribosomal RNA gene, partial sequence	0.075
653	XM_043252	Homo sapiens PCAF associated factor 65 alpha (PAF65A), mRNA	2E-33

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	Ŷ	Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
1 1		Streptococcus pneumoniae type 19C Cps19CR (cps19CR) gene,	l
		partial cds; putative oligosaccharide repeat unit transporter (cps19CJ),	
	ATTIONIC	UDP-N-acetyl glucosamine-2-epimerase (cps19CK), and putative	١
654	AF105116	glucosyl transferase (cps19CS) genes, complete cds; and gluc>	1.9
655	AF181720	Homo sapiens RU2AS (RU2) gene, complete cds, and RU2S (RU2) gene, partial cds	
055	AF 101/20	Human DNA sequence from clone RP11-269L6 on chromosome X,	0.23
656	AL392164	complete sequence [Homo sapiens]	0.0003
050	ALIJIZIOT	complete sequence promo sapiens	0,0003
657	AF189720	Oryctolagus cuniculus chloride channel (CFTR) mRNA, complete cds	0.18
	12.107720	organization desirate distance (ex 110) metric complete cas	0.10
658	AF077821	Canis familiaris inducible nitric oxide synthase mRNA, complete cds	0.077
659	NC 001327	Ascaris suum mitochondrion, complete genome	0.074
660	XM 003164	Homo sapiens KIAA0218 gene product (KIAA0218), mRNA	5.8
661	M22207	T.gratilla 217g gene encoding 217g protein, complete cds	0.071
		Mus musculus, Similar to cofactor required for Sp1 transcriptional	
ĺĺ		activation, subunit 6 (77kD), clone IMAGE:3590309, mRNA, partial	
662	BC004062	cds	0.22
663	AF007132	Homo sapiens clone 23551 mRNA sequence	0.023
		Homo sapiens hypothetical protein from clone 24796 (LOC57146),	
666	XM_007847	mRNA	5E-32
		Drosophila melanogaster genomic scaffold 142000013385531,	
667	AE002755	complete sequence	5.8
		Clostridium botulinum unidentified protein P-48 gene, complete cds	
668	U70780	and neurotoxin binding protein gene, partial cds	0.075
669	373.6 000101	Homo sapiens similar to KIAA0773 gene product (H. sapiens)	_
009	XM_029131	(LOC90115), mRNA	2
		Homo sapiens KIAA0001 gene product; putative G-protein-coupled receptor; G protein coupled receptor for UDP-glucose (KIAA0001).	
670	XM 003091	mRNA	1.7
0,0	7441_003071	Carex pauciflora tRNA-Thr, partial sequence; trnT-trnL intergenic	1.7
		spacer, tRNA-Leu, and trnL-trnF intergenic spacer, complete	
		sequence, and tRNA-Phe, partial sequence; chloroplast genes for	
671	AF284881	chloroplast products	1.6
672	NC_001381	Plasmid pAL5000, complete sequence	6.1
673	AF081201	Arabidopsis thaliana villin 1 (VLN1) mRNA, complete cds	0.21
		Homo sapiens cDNA FLJ13612 fis, clone PLACE1010833, weakly	
674	AK023674	similar to CALTRACTIN	0.54
675	X60097	Streptococcus sp. (group C) emm gene for M protein	0.063
676	AF026087	Schizosaccharomyces pombe Sat1 gene, partial cds	0.22
677	X00117	Galago Alu repeat type I, GAL9	0,0001
		Rattus norvegicus Sacm21/RT1-A intergenic region, haplotype RT1n	
678	AJ276629	and partial RT1-A gene for MHC Class I antigen	1,9
		Mus musculus adult male corpus striatum cDNA, RIKEN full-length	
679	AK021058	enriched library, clone: C030009O12, full insert sequence	0.23

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		Table 3A. Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Mus musculus, Similar to Interleukin 1 alpha, clone MGC:5780	
680	BC003727	IMAGE:3599550, mRNA, complete cds	1.9
		Homo sapiens similar to ring finger protein 23; RING-B box-coiled	
681	XM_048261	coil-B30.2 (M. musculus) (LOC92947), mRNA	5E-18
682	XM_038154	Homo sapiens KIAA1016 protein (KIAA1016), mRNA	0.23
		Homo sapiens similar to hypothetical protein MGC10940 (H. sapiens)	
683	XM_049918	(LOC93230), mRNA	0
684	X14727	Sheep beta-B globin gene	0.68
685	AB052187	Macaca fascicularis brain cDNA, clone:QnpA-12170	0.024
		Mus musculus adult male lung cDNA, RIKEN full-length enriched	
686	AK004783	library, clone:1200015E14, full insert sequence	5.8
		Drosophila melanogaster genomic scaffold 142000013385566.	
687	AE002799	complete sequence	0.003
		Corymbia calophylla tRNA-Leu and tRNA-Phe genes, partial	
688	AF190395	sequence; chloroplast genes for chloroplast products	1.9
689	U57896	Candida albicans cytochrome C gene (CYC1) gene, complete cds	0.22
690	AE006710	Sulfolobus solfataricus section 69 of 272 of the complete genome	2
		Human DNA sequence from clone RP11-269L6 on chromosome X,	
691	AL392164	complete sequence [Homo sapiens]	0,0003
	12272101	Homo sapiens immunoglobulin lambda gene locus DNA, clone:47H9	
692	D87001 `	downstream contig	0.008
693	Y13853	Drosophila erecta clone Erl inactive Bari-1 family transposon	0.16
0,,,	113055	Oryctolagus cuniculus HAC4 mRNA for hyperpolarization activated	0.10
694	AB022927	cation channel, complete cds	0,66
024	THOULETE	Araraquara virus medium RNA segment, G1/G2 glycoprotein	0.00
695	AF307327	precursor gene, partial cds	2
0,5	74 307327	Staphylococcus epidermidis strain SR1 clone step.1002g02 genomic	
696	AF269369	sequence	0.57
0,00	Al-209309	sequence	0.57
697	XM 043252	Homo sapiens PCAF associated factor 65 alpha (PAF65A), mRNA	2E-33
057	AIVI_043232	Homo sapiens I cAl associated factor of aipita (1 At OA), interval Homo sapiens lethal giant larvae (Drosophila) homolog 1 (LLGL1),	212-33
698	XM 008640	mRNA	0.22
699	AC024829	Caenorhabditis elegans cosmid Y55F3BM, complete sequence	0.023
099	AC024027	Dictyostelium discoideum clathrin heavy chain (chcA) mRNA,	0.023
700	M83660	complete cds	1.9
701	X95481	S.cerevisiae ARC1 gene	1.9
701	A73401	Homo sapiens chromosomal breakpoint fragment derivative 4,	1.7
702	AJ408897	AF4/MLL fusion, patient UPN006	4.9
702	M3408897	Molgula oculata p68 RNA helicase (bobcat) and transcription factor	4.9
703	AF110007	(manx) genes, complete cds	0.68
/03	Ariiou/	(many) gones, complete cus	0.06
		Timon DNA common Com June DDA 90711 on observe Co	
704	AL133339	Human DNA sequence from clone RP4-807J1 on chromosome 20.	2
704	U10116	Contains a GSS and a CpG island, complete sequence [Homo sapiens] Human superoxide dismutase (SOD3) gene, complete cds	0.23
706			
/06	AB037771	Homo sapiens mRNA for KIAA1350 protein, partial cds	0

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		Table 3A Nearcst Neighbor (BlastN vs. Genbank)	,
SEQ ID	A COURTON	Drace.	
NO	ACCESSN	DESCRIP	P VALUE
707	AF023840	Homo sapiens natural killer group protein 2-A (NKG2-A) gene, complete cds	0,68
		Tegeticula maculata extranea cytochrome oxidase subunit I,	
		cytochrome oxidase subunit II genes, mitochondrial genes encoding	ĺ
		mitochondrial proteins, partial cds, and tRNA-Leu gene,	
708	U49023	mitochondrial gene	0.001
700	77.4.044	Homo sapiens similar to death receptor 6 (H. sapiens) (LOC92385),	
709	XM_044768	mRNA	e-126
710	D50006	Human DNA for alpha-platelet-derived growth factor receptor, exon 6-	٠
/10	D30000	III	2.1
711	Y14524	Tetrahymena thermophila tRNA-Tyr gene, exons 1 and 2, clone pTetY2	
/11	1 14324	Human DNA sequence from clone RP13-237P14 on chromosome X.	0,057
712	AL592159	complete sequence [Homo sapiens]	0.52
713	U88825	HIV-1 isolate 92NG003 from Nigeria complete genome	0.52 3.1
713	000023	Blastocrithidia culicis ATCC 30268 kinetoplast (MURF1) and	3.1
		apocytochrome B (cytB) genes, partial cds, and ATPase subunit 6	1
714	U05813	(MURF4) gene, complete pre-RNA-edited DNA sequence	0.23
715	AF048990	Homo sapiens MutS homolog 5 (MSH5) gene, exons 11 and 12	0.0009
	111010000	Adoncholaimus thalassophygas isolate Sou39 large subunit ribosomal	0.0002
		RNA gene, partial sequence; mitochondrial gene for mitochondrial	
716	AF317082	product	0,0008
720	12017002	Mus musculus 10, 11 days embryo cDNA, RIKEN full-length	0,0000
717	AK013269	enriched library, clone:2810439K08, full insert sequence	0,024
		Physarum polycephalum cytochrome oxidase subunit 1 mRNA.	-1,0-1
718	L14769	complete cds; mitochondrial gene for mitochondrial product	1.6
720	U45453	Xenopus laevis transcription factor xGATA-4 mRNA, complete cds	0.61
721	AF090432	Danio rerio serrateB mRNA, complete cds	0,075
722	M62504	Spiroplasma citri fibril protein gene, complete cds	0.024
723	AK000939	Homo sapiens cDNA FLJ10077 fis, clone HEMBA1001864	0
724	XM_032587	Homo sapiens 2127 (KIAA1754), mRNA	0.22
		Sambucus nigra ribosome inactivating protein precursor mRNA,	
726	AF012899	complete cds	0.000001
		RAR-gamma 2.2=retinoic acid receptor gamma isoform 2.2	
		{alternatively spliced} [Xenopus laevis, embryos, stage 24, mRNA,	
727	S82175	2154 nt]	0,008
		Mus musculus adult male kidney cDNA, RIKEN full-length enriched	
728	AK002856	library, clone:0610040A09, full insert sequence	0.44
		Oryctolagus cuniculus indolethylamine N-methyltransferase (INMT)	l
729	AF077828	gene, exon 3 and complete cds	1.4
	m.coo.c.c.	Mus musculus, Flt3 interacting zinc finger protein 1, clone	
730	BC006633	MGC:7201 IMAGE:3482191, mRNA, complete cds	1.3
731	AK024670 Z35823	Homo sapiens cDNA: FLJ21017 fis, clone CAE05907	0.5
732 733	Z35823 Z48636	S.cerevisiae chromosome II reading frame ORF YBL062w	1.1
133	Z/46030	C.novyi gene for alpha-toxin	0.065

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3734 BC008380 Homo sspiens, clone IMAGE:4179986, mRNA, partial cds	P VALUE 0.71 0.11 0.95 0.48 0.5
	0.71 0.11 0.95 0.48 0.5
1935	0.11 0.95 0.48 0.5 1.6 0.034
Anolis roquet cytochrome b gene, partial cds; mitochondrial gene for mitochondrial product AF142372 Anolis roquet cytochrome b gene, partial cds; mitochondrial gene for mitochondrial product Britonesspiens solute carrier family 6 (neurotransmitter transporter), memory and propher of (SLC6AH), mRNA Arabidopsis thaliana clone AtDP5 isopentenyl prophosphate-dimethyliallyl prophosphate isomerase (ipiAt2) mRNA, partial cds AF188067 ANO X90383 Athaliana DNA for Y13 gene Vibrio cholerae chromosome I, section 217 of 251 of the complete chromosome Calothrix partielina clone 102-2A 16S-23S internal transcribed spacer, complete sequence; and tRNA-Ila and tRNA-Ala genes, complete sequence; and tRNA-Ila and tRNA-Ala genes, complete sequence; and tRNA-Ila and tRNA-Ala genes, complete sequence; and tRNA-Ila gene, partial cds XM_001795 Homo sapiens RING3 protein (KIAA0876), mRNA Pantellella bicolor cytochrome oxidase subumi I gene, partial cds, matchondrial gene for mitochondrial product Homo sapiens clone RIOMS NAIL1418 chromosome 15 paralogous sequence AF262633 AF262633 AF262633 AF262633 AF262634 Mis musculus TFFI/pS2 gene for Trefoil Factor I/pS2, exons 1-3 AF0524 AL031634 Calorababilis elegans cosmul Y32BIZC, complete sequence	0.95 0.48 0.5 1.6 0.034
373	0.48 0.5 1.6 0.034
	0.48 0.5 1.6 0.034
Homo sepiens solute carrier family 6 (neurotransmitter transporter), member 14 (SLC6A14), mRNA. Arabidopsis thaliana clone AIDP5 isopentenyl pyrophosphate cimethyllallyl pyrophosphate isomerase (ipiAt2) 739	0.5 1.6 0.034
38	1.6 0.034
Arabidopsis thaliana clone AIDP5 isopentenyl prophosphate: dimethyliallyl prophosphate isomerase (ipiAt2) mRNA, partial cds Arabidopsis thaliana DNA for Yi3 gene Vibiro cholerae chromosome I, section 217 of 251 of the complete chromosome Calothrix partielina clone 102-2A 16S-23S internal transcribed spacer, complete sequence; and tRNA-Ila and tRNA-Ala genes, complete sequence AF236642 AF236642 AF276079 Homo sapiens RING3 protein (RING3) gene, partial cds SM_035624 Homo sapiens RING3 protein (RING3) gene, partial cds XM_001795 Homo sapiens RING3 protein (RING3) gene, partial cds XM_001795 Homo sapiens RING3 protein (RIAA0876), mRNA Partellella bicolor cytochrome oxidase subumit I gene, partial cds, mitochondrial gene for mitochondrial product Homo sapiens clone RIOMS NAIL1418 chromosome 15 paralogous AF262633 Soppiens clone RIOMS NAIL1418 chromosome 15 paralogous AF2636375 Homo sapiens clone RIOMS NAIL1418 chromosome 15 paralogous AF26363175 Homo sapiens fulle might insert cDNA clone ZD68B12 Mis musculus TFFI/pS2 gene for Trefoil Factor I/pS2, exons 1-3 AF063681 Mus musculus type XIII collagen (coll3a1) gene, exons 22-27 AF391634 Calonadoria cosmid Y32B12C, complete sequence AF301634 Calonadoria cosmid	1.6 0.034
	0.034
AF188067 mRNA, partial cds	0.034
740	0.034
Vibrio cholerae chromosome I, section 217 of 251 of the complete chromosome Calothrix parietina clone 102-2A 16S-23S internal transcribed spacer, complete sequence; and tRNA-IIa and tRNA-AIa genes, complete sequence; and tRNA-IIa and tRNA-AIa genes, complete sequence; and tRNA-IIa and tRNA-AIa genes, complete sequence; and tRNA-IIIa and tRNA-AIa genes, complete sequence (STA-IIIIa) and tRNA-AIa genes, complete sequence (RIA-AIB (STA-IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	
741 AB004309 chromosome Calothrix parietina clone 102-2A 16S-23S internal transcribed spacer, complete sequence; and fRNA-lla and fRNA-Ala genes, complete sequence 742 AF236642 sequence 744 AF107699 Homo sapiens RING3 protein (RING3) gene, partial cds 745 XM_035624 Homo sapiens KIAA0876 protein (KIAA0876), mRNA 746 XM_001795 Homo sapiens ISIAA0876 protein (KIAA0876), mRNA 747 AF395180 mitochondrial gene for mitochondrial product 748 AF262633 sequence variant, genomic sequence 749 AF086375 Homo sapiens tolne RIOMS NAIL1418 chromosome 15 paralogous sequence sepiens fulle Inght insert cDNA clone ZD68B12 750 AI271002 Mus musculus TFF1/pS2 gene for Trefoil Factor 1/pS2, exons 1-3 751 AZ036341 Mus musculus cosmid Y32B17C, complete sequence	0.18
Calothrix parietina clone 102-2A 16S-23S internal transcribed spacer, complete sequence, and tRNA-lle and tRNA-Ala genes, complete sequence and tRNA-lle and tRNA-Ala genes, complete sequence RNA-lle and tRNA-Ala genes, complete sequence RNA-lle and tRNA-Ala genes, complete sequence RNA-genes, complete sequence RNA-genes, complete sequence RNA-genes, complete sequence Roman Sequence Roman Sequence Roman Sequence Name Sequence Sequence Sequence RNA-genes	0.18
complete sequence; and tRNA-Ile and tRNA-Ala genes, complete sequence	
742 AF236642 sequence 744 AF107699 Homo sapiens RING3 protein (RING3) gene, partial cds 745 XM 035624 Homo sapiens KIAA0876 protein (KIAA0876), mRNA 746 XM_001795 Homo sapiens Isamin B receptor (LBR), mRNA 747 AF395180 mitochondrial gene for mitochondrial product 748 AF262633 sequence variant, genomic sequence 749 AF086375 Homo sapiens full length insert cDNA done ZD68B12 750 AJ271002 Mus musculus TFF1/pS2 gene for Trefoil Factor Ip/S2, exons 1-3 751 AF063681 Mus musculus type XIII collagen (coll3a1) gene, exons 22-27 752 AL031634 Caenorhabditis elegans cosmid Y32B12C, complete sequence	
744 AF107699 Homo sapiens RING3 protein (RING3) gene, partial cds 745 XM_035624 Homo sepiens RINA0876 protein (KIAA0876), mRNA 746 XM_001795 Homo sapiens afmin E receptor (LIBN, mRNA 747 AF395180 Panteliella bicolor cytochrome oxidase subunit I gene, partial cds; mitochondrial gene for mitochondrial product 748 AF262633 mitochondrial gene for mitochondrial product 749 AF086375 Homo sepiens full length insert cDNA done ZD68B12 750 AJ271002 Mus musculus TFF1/pS2 gene for Trefail Factor1/pS2, exons 1-3 751 AF063681 Mus musculus type XIII collager (coll 3a1) gene, exons 22-27 752 AL031634 Casendrabditis elegans cosmid 732B12C, complete sequence	
745 XM_035624 Homo sepiens KIAA0876 protein (KIAA0876), mRNA 746 XM_001795 Homo sepiens lamin B receptor (LBR), mRNA 747 AF395180 mitochondrial gene for mitochondrial product 748 AF262633 sepiens clone NIGMS NAIL1418 chromosome 15 paralogous 749 AF068375 Homo sepiens tolne NIGMS NAIL1418 chromosome 15 paralogous 750 AZ71002 Mins musculus TFF1/pS2 gene for Trefoil Factor I/pS2, exons 1-3 751 AZ036381 Mus musculus type XIII collagen (coll 3a1) gene, exons 22-27 752 AL031634 Casenorhabditis elegans cosmid 732B12C, complete sequence	1.1
746 XM_001795 Homo sapiens Iamin B receptor (LBR), mRNA 747 AF395180 mitochondrial gene for mitochondrial product 748 AF262633 mitochondrial gene for mitochondrial product 749 AF086375 Homo sapiens clone NIGMS NA11418 chromosome 15 paralogous sequence variant, genomic sequence 750 AJ271002 Mus musculus TFFI/pS2 gene for Treful Factor I/pS2, exons 1-3 751 AF063681 Mus musculus type XIII collagen (coll3a1) gene, exons 22-27 752 AL031634 Caenorhabditis elegans cosmid Y32B12C, complete sequence	0.027
Parteliella bicolor cytochrome oxidase subunit I gene, partial cds; mitochondrial gene for mitochondrial product Homo sapiens clone NIGMS NAIL18 chromosome 15 paralogous sequence variant, genomic sequence AF262633 Homo sapiens full english insert cDNA clone ZD68B12 750 AJ271002 Mus musculus TFFI/pS2 gene for Trefoil Factor I/pS2, exons 1-3 AF063631 Mus musculus type XIII collagen (coll 3a1) gene, exons 22-27 752 AL031634 Casenrhadditis elegans cosmid Y32B1C2, complete sequence AF262632 AF263634 AF2636344 AF263644 AF263	3.4
747 AF395180 mitochondrial gene for mitochondrial product Homo sapiens (olne NIGMS NA11418 chromosome 15 paralogous sequence variant, genomic sequence 749 AF086375 Homo sapiens full length insert cDNA clone ZD68B12 750 AJ271002 Mus musculus TFFI/pS2 gene for Trefoil Factor I/pS2, exons 1-3 751 AF063681 Mus musculus type XIII collagen (coll3a1) gene, exons 22-27 752 AL031634 Caenorhabditis elegans cosmid Y32B12C, complete sequence	0.15
Homo sapiens clone NIGMS NA11418 chromosome 15 paralogous sequence variant, genomic sequence AF086375 Homo sapiens full length insert cDNA clone ZD68B12	
748 AF262633 sequence variant, genomic sequence 749 AF086375 Homo spiens full length insert cDNA clone ZD68B12 750 AJ27102 Mns musculus TFF1/pS2 gene for Trefoil Factor1/pS2, exons 1-3 751 AF063681 Mus musculus type XIII collagen (coll3a1) gene, exons 22-27 752 AL031634 Caenorhabditis elegans cosmid Y32B12C, complete sequence	0.093
749 AF086375 Homo sepiens full length insert cDNA clone ZD68B12 750 AJ271002 Mus musculus TFF1/pS2 gene for Trefoil Factor 1/pS2, exons 1-3 751 AF063681 Mus musculus type XIII collagen (coll3a1) gene, exons 22-27 752 AL031634 Caenorhabditis elegans cosmid Y32B12C, complete sequence	
750 AJ271002 Mns musculus TFFL/pS2 gene for Trefoil Factor L/pS2, exons 1-3 751 AF063681 Mus musculus type XIII ollagen (coll13a) gene, exons 22-27 752 AL031634 Caenorhabditis elegans cosmid Y32B12C, complete sequence	0.27
751 AF063681 Mus musculus type XIII collagen (coll3a1) gene, exons 22-27 752 AL031634 Caenorhabditis elegans cosmid Y32B12C, complete sequence	0.000003
752 AL031634 Caenorhabditis elegans cosmid Y32B12C, complete sequence	0.53
	0.053
	0.055
Homo sapiens genomic DNA, chromosome 21q22.2, PCR fragment	0.003
754 AP000693 from BAC clone:KB739C11, CBR1-HLCS region	0.022
755 AK021535 Homo sapiens cDNA FLJ11473 fis, clone HEMBA1001712	0.022
Mus musculus B lymphocyte induced maturation protein 1 (Prdm1)	0.004
756 AF305539 gene, exon 8 and complete cds	4E-12
730 At 303337 gene, exon 6 and compact cus	715-12
757 J05493 S.pombe no message in thiamine protein (nmt1) gene, complete eds	0.55
Helicobacter pylori, strain J99 section 41 of 132 of the complete	
758 AE001480 genome	0.51
Bos taurus matrix metalloproteinase 13 (MMP13) mRNA, complete	
759 AF072685 cds	0.072
760 XM 038408 Homo sapiens hypothetical protein FLJ11331 (FLJ11331), mRNA	0.6
Homo sapiens hypothetical gene supported by AF037219	
761 XM 041148 (LOC91856), mRNA	1.7
Haemulon plumicri Hpl224 NADH dehydrogenase I (ND-1) gene,	
762 AF234897 complete cds; mitochondrial gene for mitochondrial product	3

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
763	NM_008399	Mus musculus integrin, alpha E, epithelial-associated (Itgae), mRNA	1.4
764	Z74898	S.cerevisiae chromosome XV reading frame ORF YOL156w	1.8
765	AF280812	Glycine max putative Hs1pro-1-like receptor mRNA, complete cds	0.59
766	AB013615	Broad bean wilt virus 2 genomic RNA, complete sequence of RNA1	0.68
	17050150	Nicotiana tabacum feedback-insensitive anthranilate synthase alpha-2	
768	AF079169	chain (ASA2) gene, promoter region and 5'UTR	5.9
		Homo sapiens genomic DNA, chromosome 6p21.3, HLA class I	
769	AB014079	region, Cosmid clone:TY1E11, complete sequence	0.007
770	XM_012336	Homo sapiens MBIP protein (MBIP), mRNA	0.0003
771	U41509	Caenorhabditis elegans cosmid W03H1, complete sequence	0.18
772	U23478	Dictyostelium discoideum phosphatidylinositol-4,5-diphosphate 3- kinase (PIK3) mRNA, partial cds	0.00002
		Arabidopsis thaliana genomic DNA, chromosome 5, genomic PCR	0,00002
773	AP000380	product GA469	0.39
		Danio rerio basic helix-loop-helix transcription factor Ndr1a (ndr1a)	
774	AF115772	mRNA, complete cds	0.2
775	XM_007458	Homo sapiens butyrate response factor 1 (EGF-response factor 1) (BRF1), mRNA	0.42
776	L31955	Human (clone 1NIB-2) normalized cDNA library sequence	1.6
		Populus kitakamiensis (P. sieboldii X P. grandidentata) homt3 gene	
777	D49711	for caffeic acid O-methyltransferase, complete cds (exon1-4)	0.046
778	L49234	Manduca sexta chitinase precursor gene, complete cds	0.05
		Ictalurus punctatus clone Icpu-UA/3 MHC class I antigen gene,	
779	AY008848	complete cds	0.015
780	AB048375	Oserya coulteriana chloroplast matK gene for maturase K, complete cds	1.9
781	XM 010323	Homo sapiens toll-like receptor 7 (TLR7), mRNA	0.21
782	X69121	P.falciparum Pfgrp mRNA for heat shock protein	0.18
702	2007121	Helicobacter pylori, strain J99 section 131 of 132 of the complete	0.10
783	AE001570	genome	0.05
784	M31756	Human t(9;22) chromosome 9 breakpoint DNA	0.77
		Staphylococcus aureus replication (rep), control of replication (cop),	
785	M37889	and resistance protein (QacC) genes, complete cds	0.2
		Plasmodium falciparum histone deacetylase (HDAC1) mRNA,	
786	AF091326	complete cds	0.19
787	Z74851	S.cerevisiae chromosome XV reading frame ORF YOL109w	0.2
		H. sapiens Ig lambda light chain variable region gene (34-34SWIIF32)	
788	Z85396	rcarranged; Ig-Light-Lambda; VLambda	0.56
789	AF075681	Hemorrhagic enteritis virus hexon protein gene, partial cds	2.1
		Mus musculus GTP binding protein 3 gene, complete cds; nuclear	
790	AY029613	gene for mitochondrial product	5.8
791	NC_001338	Sulfolobus virus 1, complete genome	0.63

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WC0214500 [flig //E /WO0214500 opc]

The complete genome The complete genome	-			
NO ACCESSN DESCRIP P VALUE			Table 3A Nearest Neighbor (BlastN vs. Genbank)	
The complete genome The complete genome	SEQ ID			
Total	NO	ACCESSN	DESCRIP	P VALUE
Total				
AF293450 Canis familiaris BPAGI-like gene sequence 0.53	792	U67563	Methanococcus jannaschii section 105 of 150 of the complete genome	0.11
AF293450 Canis familiaris BPAGI-like gene sequence 0.53				
	794	U67549	Methanococcus jannaschii section 91 of 150 of the complete genome	0.21
Homo sapiens, Similar to hypothetical protein RL10842, clone			Canis familiaris BPAG1-like gene sequence	0.53
Post	797	XM_029942	Homo sapiens KIAA0877 protein (KIAA0877), mRNA	1.4
NM_008399 Mus musculus integrin, alpha E, epithelial-associated (Itgae), mRNA 0.24			Homo sapiens, Similar to hypothetical protein FLJ10842, clone	
Methanococcus jannaschii small extra-chromosomal element, complete sequence 2 2 298532 Spombe chromosome I cosmid c1B1 0.66	798	BC009775	MGC:13398 IMAGE:4092662, mRNA, complete cds	0.061
Methanococcus jannaschii small extra-chromosomal element, complete sequence 2 2 298532 Spombe chromosome I cosmid c1B1 0.66				
NC_001733 complete sequence 2	799	NM_008399		0.24
Sport Spor				
Human DNA sequence from clone RP3-323A24 on chromosome 4. Contains GSSs and a putative CpG island, complete sequence [Homo Contains GSSs and a putative CpG island, complete sequence [Homo Sapiens] 0,003				2
AL158132 Saplens	801	Z98532		0.66
AL158132 sapiens 0.003				
Homo supiens hypothetical protein from clone 24796 (LOC57146), mRNA 5E-32				
SM_007847 mRNA 5E-32 mRNA	802	AL158132		0.003
Mandarina mandarina from Antjima population 16S ribosomal RNA gene, mitochondrial gene encoding mitochondrial product, partial sequence 0.086				
	803	XM_007847	mRNA	5E-32
Mandarina mandarina from Anijima population 16S ribosomal RNA gene, mitochondrial gene encoding mitochondrial product, partial sequence 0.086				
AF098697 sequence 0.086	805	XM_003799		0.71
AF007509 Streptococcus pneumoniae section 192 of 194 of the complete genome 1.6	1			
AE007509 Streptococcus pneumoniae soction 192 of 194 of the complete genome 1.6		.=		
Arabidopsis thaliana unknown protein (T20O10_100/AT3g63000) 0.2	806	AF098697	sequence	0.086
Arabidopsis thaliana unknown protein (T20O10_100/AT3g63000) 0.2	007	A E COST COO	St	
AP035064 mRNA, complete cds 0.2	807	AE00/509		1.0
AE000595 Helicobacter pylori 26695 section 73 of 134 of the complete genome 0.046	909	A3702.506.4		0.2
Single Scerevisiae chromosome XV reading frame ORF YOR185c 0.11	808	A1033004	Inrava, complete cus	0.2
Single Scerevisiae chromosome XV reading frame ORF YOR185c 0.11	900	AE000505	Welicohagter puleri 26605 section 73 of 134 of the complete genome	0.046
M30825 Drosophila melanogaster ovarian tumor protein isoforms (otu) gene, complete cds, alternatively spliced 0.11				
M30825 complete cds, alternatively spliced 0.11	010	2/3093		0.11
S12	911	M30925		0.11
Homo sapiens, Similar to RIKEN cDNA 5830420C20 gene, clone 1.7				
BC005104 MAGE:3633379, mRNA, partial cds 1.7		117007		0.005
S14 XM_029941 Homo sapiens cadherin-like 22 (CDH22), mRNA 0.0006	813	BC005104		17
815 NM_003246 Homo sapiens thrombospondin 1 (THBS1), mRNA 0.23				
Canis familiaris junctional sarcoplasmic reticulum protein mRNA, complete cds 0.21	815			0.23
816 U38414 complete cds 0.21				
Homo sapiens chromodomain helicase DNA binding protein 1 (CHD1), mRNA Cryptococcus neoformans var. grubii strain H99 urease (URE1) gene, complete cds 1.7	816	U38414		0.21
S17 XM_004000 (CHD1), mRNA 0.63				
Cryptococcus neoformans var. grubii strain H99 urease (URE1) gene, complete cds 1.7	817	XM 004000		0.63
818 AF006062 complete cds 1.7				
	818	AF006062		1.7
010 774 040000 77				
819 XM_018303 Homo sapiens hypothetical protein FLJ11565 (FLJ11565), mRNA 0.07	819	XM_018303	Homo sapiens hypothetical protein FLJ11565 (FLJ11565), mRNA	0.07

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
820	AE001164	Borrelia burgdorferi (section 50 of 70) of the complete genome	1.6
821	AF022969	Caenorhabditis elegans cosmid C29G2	0.2
822	AK021462	Homo sapiens cDNA FLJ11400 fis, clone HEMBA1000673	1.6
823	Y10908	B.cereus cysA and wapA genes	0.0003
824	U36892	Triticum aestivum clone AC29 microsatellite sequence	3.4
825	AF258615	Dictyostelium discoideum LagC protein (lagC) gene, complete cds	0.011
826	Z99763	Flaveria pringlei gdcsH gene	0.023
827	X71543	B.taurus microsatellite sequence INRA118	7E-15
		Arabidopsis thaliana putative 60S ribosomal protein (T6C23.18)	
828	AF349526	mRNA, complete cds	1.7
		Homo sapiens immunoglobulin lambda gene locus DNA, clone:47H9	
829	D87001	downstream contig	0.007
830	XM_006470	Homo sapiens radixin (RDX), mRNA	2.4
831	AF191609	Homo sapiens filamin (FLNB) gene, exon 21	e-140
1		Homo sapiens nuclear inhibitor of protein phosphatase-1 (PPP1R8)	
832	AF064754	gene, exon 4	5.5
833	NC_001438	Bean golden mosaic virus A component DNA, complete sequence	0.57
834	D14567	Penicillium urticae mitochondrial I-rRNA (large rRNA) gene and its flanking region	0.23
		Homo sapiens genomic DNA, chromosome 21q22.1, D21S226-AML	
835	AP000301	region, clone:D16B8, complete sequence	0.5
836	NM_014147	Homo sapiens HSPC047 protein (HSPC047), mRNA	1E-90
837	AF296100	Porcine teschovirus strain DS 562/91 polyprotein gene, complete cds	1.3
		Mus musculus adult male spinal cord cDNA, RIKEN full-length	
838	AK020740	enriched library, clone: A330106M24, full insert sequence	1.6
839	XM_045585	Homo sapiens hypothetical protein FLJ11085 (KIAA1434), mRNA	3E-16
840	AF115104	Homo sapiens glutaredoxin (GLRX) gene, exon 1	1.3
841	AF094673	Mus musculus Sos1 gene, intron	3E-26
842	AK025101	Homo sapiens cDNA: FLJ21448 fis, clone COL04473	5E-93
843	M85276	Homo sapiens NKG5 gene, complete cds	0.039
844	AK022390	Homo sapiens cDNA FLJ12328 fis, clone MAMMA1002145	0.018
845	AF269767	Staphylococcus epidermidis strain SR1 clone step.1023b06 genomic sequence	0.47
545	111 237707	Homo sapiens minichromosome maintenance deficient (S. cerevisiae)	0.47
846	NM_005916	7 (MCM7), mRNA	4.5
		Rattus norvegicus stromal cell-derived factor 1 (SDF-1) mRNA,	
847	AF209976	complete cds	1.3
848	NM_017078	Rattus norvegicus Acetylcholine receptor alpha 5 (Chrna5), mRNA	0.95
		Staphylococcus epidermidis strain SR1 clone step.1002a12 genomic	
849	AF269324	sequence	0.16

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
	THE GENERAL	Mus musculus adult male stomach cDNA, RIKEN full-length	X VILLOUI
850	AK008669	enriched library, clone:2210008F06, full insert sequence	4.3
851	XM 044217	Homo sapiens Ran binding protein 11 (LOC51194), mRNA	0.44
		Homo sapiens CDC5 (cell division cycle 5, S. pombe, homolog)-like	
852	XM_011510	(CDC5L), mRNA	0.056
853	AF068574	Arabidopsis thaliana ferulate-5-hydroxylase (F5H) gene, complete cds	0.042
		Drosophila melanogaster genomic scaffold 142000013386025,	
854	AE003072	complete sequence	1.4
855	AF159462	Escherichia coli EHEC factor for adherence (efa1) gene, complete cds	0.42
	111107111	Rattus norvegicus GABAB1 receptor (GABABR1) gene, exons 10	0.12
856	AF110797	through 20 and alternatively spliced products, complete cds	1.6
	111 110/7/	Homo sapiens cGMP-specific phosphodiesterase type 5A (PDE5A)	1.0
857	AF319172	gene, partial cds	0.6
	400	Human chromosome 10 duplicated adrenoleukodystrophy (ALD)	
858	U90299	gene, 3' breakpoint sequence	0.0000003
		Homo sapiens mRNA; cDNA DKFZp566M043 (from clone	
859	AL050065	DKFZp566M043)	0.006
		Haemophilus influenzae Rd section 117 of 163 of the complete	
860	U32802	genome	1.4
861	AK001127	Homo sapiens cDNA FLJ10265 fis, clone HEMBB1001014	0.17
862	AB042995	Plecoglossus altivelis DNA, microsatellite, clone:Pal-3*	0.45
863	NM_004668	Homo sapiens maltase-glucoamylase (alpha-glucosidase) (MGAM), mRNA	0.001
		Rattus norvegicus neuronal potassium channel alpha subunit (Kv8.1),	
864	NM_021697	mRNA	0.19
865	Z28173	S.cerevisiae chromosome XI reading frame ORF YKL173w	0.035
866	XM_051783	Homo sapiens hypothetical protein FLJ11712 (FLJ11712), mRNA	1.1
867	X92498	M.musculus fkh-6 gene	0.99
0.0	70 5 041000	Homo sapiens similar to phosphodiesterase 4D interacting protein	
868	XM_041339	(myomegalin) (H. sapiens) (LOC91892), mRNA	1.5
0.00		Human DNA sequence from clone RP11-466P12 on chromosome 6,	
869	AL590401	complete sequence [Homo sapiens]	0.16
]		Thomas DNA someone from also I I 22NG02 85U10 on also on a	
870	Z68758	Human DNA sequence from clone LL22NC03-85E10 on chromosome 22 Contais GSSs, complete sequence [Homo sapiens]	0.021
670	200730	22 Contais G558, Complete sequence [riomo sapiens]	0.021
871	Z34916	O.aries DNA for natural resistance associated macrophage protein	4.6
		Mus musculus serine protease inhibitor-2 related sequence 1 (Spi2-	
872	NM_009253	rs1), mRNA	1.5
9770	13700 1000	Oryza sativa microsatellite MRG6608 containing (TGTA)X6,	
873 874	AY024283	genomic sequence	3.8
874	Z36108	S. cerevisiae chromosome II reading frame ORF YBR239c Homo sapiens hypothetical protein (LOC51320), mRNA	0.14
0/0	XM_008791	priomo sapiens nypotneticai protein (LOC>1320), mKNA	0.14

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SEO ID		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID	ACCESSN	DESCRIP	P VALUE
876	X89383	R. norvegicus mRNA for SNF1-related kinase	0.18
	1107505	Homo sapiens uncharacterized hypothalamus protein HT010 (HT010),	0.10
877	XM_002378	mRNA	0
879	XM_047537	Homo sapiens DKFZP434C153 protein (DKFZP434C153), mRNA	0.003
	3242_017337	Homo sapiens checkpoint with forkhead and ring finger domains	0.003
880	XM_006700	(CHFR), mRNA	4E-27
881	NC_002756	Mertensiella luschani mitochondrion, complete genome	2,1
		Human immunodeficiency virus type 1, viral sample LC01.DA11, V3	
882	M90915	region	0.019
883	AB011534	Rattus norvegicus mRNA for MEGF8, partial cds	0.19
885	AF364468	Simian-Human immunodeficiency virus clone 201/26w 1 envelope glycoprotein (env) gene, partial cds	1.4
		Homo sapiens cerberus 1 (Xenopus laevis) homolog (cysteine knot	
886	XM_005320	superfamily) (CER1), mRNA	0.63
887	AC079875	Homo sapiens clone CTD-2333B13, complete sequence	0.024
888	NM_032394	Mus musculus myosin VIIb (Myo7b), mRNA	6.6
889	AF209448	Borrelia hermsii plasmid cp32-like EcoRI-cut fragment #3	1.8
890	AK008242	Mus musculus adult male small intestine cDNA, RIKEN full-length enriched library, clone:2010015A19, full insert sequence	0.065
		Homo sapiens clone 16 ribosomal RNA intergenic spacer region,	
891	AF305851	partial sequence	2
892	XM_047604	Homo sapiens coproporphyrinogen oxidase (coproporphyria, harderoporphyria) (CPO), mRNA	0.22
893	AF211133	Carsonella ruddii natural-host Ctenarytaina eucalypti tryptophamyl- tRNA synthetase (trpS) gene, partial cds; 16S ribosomal RNA, 23S ribosomal RNA, and 5S ribosomal RNA genes, complete sequence; transaldolase (tal) and alkly lyhdyroperoxide reductase smal>	0.2
894	AY047513	Drosophila melanogaster GH01880 full length cDNA	8.8
	111047515	Human DNA sequence from clone RP3-323A24 on chromosome 4.	0.0
		Contains GSSs and a putative CpG island, complete sequence [Homo	
896	AL158132	sapiens]	0.0001
897	Z21978	B.napus Myr2.Bn1 gene encoding myrosinase, thioglucoside glucohydrolase	1.4
898	X95904	L.esculentum mRNA for 14-3-3 protein, TFT6	0.81
899	Z48930	C.caldarium mitochondrial genes (10kb)	0.81
900	XM 038535	Homo sapiens KIAA1074 protein (KIAA1074), mRNA	0.27
901	AF016898	Homo sapiens B-ATF gene, complete cds	0.002
-701	AI 010070	Homo sapiens similar to PRO0097 protein (H. sapiens) (LOC93544),	0.002
902	XM 051951	mRNA	0.003
902	X89383	R.norvegicus mRNA for SNF1-related kinase	0.003
903	A07303	Danio rerio bone morphogenetic protein 4 precursor (BMP4) gene,	0.23
904	AF056336	complete cds	0.002
905	U34339	Protopterus acthiopicus 28S ribosomal RNA gene, partial sequence	1E-09
		249	

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Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE Homo sapiens hypothetical gene supported by AK025102 906 XM 041974 (LOC91974), mRNA 0.06 Penicillium urticae mitochondrial I-rRNA (large rRNA) gene and its 907 D14567 flanking region 0.26 908 AL035632 Drosophila melanogaster BAC clone BACN32G11 0.52 Drosophila melanogaster mitotic kinesin-like motor protein CENP-909 AF220354 ana (cana) mRNA, complete cds 0.86 910 U67484 Methanococcus jannaschii section 26 of 150 of the complete genome 26 911 Y13853 Drosophila erecta clone Er1 inactive Bari-1 family transposon 0.17 Homo sapiens mRNA: cDNA DKFZp586A2324 (from clone 912 AL137643 DKFZp586A2324) 1.3 Streptococcus pyogenes M1 GAS strain SF370, section 147 of 167 of AE006618 913 the complete genome 1.4 914 Z46663 M.musculus DNA for growth hormone gene and promoter 0.005 AF197225 Emericella nidulans TUPA (tupA) gene, complete cds 915 0.65 916 L20594 Human annexin III (ANX3) gene, exon 4 0.028 Clostridium haemolyticum genes for 16S rRNA, tRNA-Ala, tRNA-Ile. 917 AB040730 23S rRNA, partial and complete sequence 0.31 Human variable number tandem repeat (VNTR) region, allele 13R4 3' 918 L10162 to collagen type II (COL2A1) gene 0.19 L35061 Bacteriophage phi-41 orfs L1-12, complete cds's, orf 13, 5' end 0.75 Mus musculus adult male cecum cDNA, RIKEN full-length enriched 920 AK018618 library, clone:9130014E20, full insert sequence 0.72 921 AC024865 Caenorhabditis elegans cosmid Y74C10AL, complete sequence 0.083 H.sapiens CpG island DNA genomic Mse1 fragment, clone 69g7, 922 Z65965 reverse read cpg69g7.rt1b 0.46 Influenza A virus (A/Goose/Guangdong/3/97(H5N1)) segment 6 923 AF364335 neuraminidase (NA) gene, complete cds 0.46 924 M83534 A.thaliana isocitrate lyase gene, 5' end 0.003 Plasmodium falciparum chromosome 2, section 27 of 73 of the 925 AE001390 complete sequence 0.0001 926 AK023864 Homo sapiens cDNA FLJ13802 fis. clone THYRO1000186 0 XM 010575 Homo sapiens LOC87729 (LOC87729), mRNA 927 1.3 928 L16771 Drosophila simulans suppressor of forked (su(f)) gene sequence 0.17 931 BC001815 Homo sapiens, clone IMAGE:2989261, mRNA 2E-33 BC008580 Homo sapiens, clone IMAGE:4179986, mRNA, partial cds 932 1.4 Human gene for 2-oxoglutarate dehydrogenase, exon 14, 15, 16, 17, 933 D32063 18 and 19 sequence 0.058 Homo sapiens hypothetical gene supported by AK026706 934 XM 028551 (LOC90063), mRNA 0.003 935 M87298 Chicken neuropeptide Y gene, exons 1-3 936 AE006100 Pasteurella multocida PM70 section 67 of 204 of the complete genome 1.5 L77040 Homo sapiens (subclone 8 c11 from P1 H22) DNA sequence 9E-14 937

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
965	AF039714	Caenorhabditis elegans cosmid T13E5	1.8
		Plasmodium falciparum 1-deoxy-D-xylulose 5-phosphate	
966	AF111813	reductoisomerase (dxr) gene, complete cds	1.3
		Escherichia coli K12 MG1655 section 315 of 400 of the complete	
967	AE000425	genome	0.002
968	AF108658	Homo sapiens p5327 mRNA, complete cds	0.57
969	X82329	A.hypogaea chi2.1 gene for chitinase (class II)	3.6
970	AK026575	Homo sapiens cDNA: FLJ22922 fis, clone KAT06722	1.5
		Bombyx mori gene for peptidoglycan recognition protein, complete	
971	AB016605	cds	0.0002
972	AB009629	Avian rotavirus RNA for VP1, complete cds	1.4
		Mus musculus 12 days embryo embryonic body between diaphragm	
		region and neck cDNA, RIKEN full-length enriched library,	
973	AK020413	clone:9430013L14, full insert sequence	0.095
		Gallus gallus transcription factor RREB-1 (rreb-1) mRNA, complete	
974	AF013754	cds	3.4
975	V01390	Trypanosoma brucei kinetoplast DNA maxicircle fragment encoding two very small ribosomal RNAs, a 12S rRNA homologous to E.coli 23S rRNA and a 9S rRNA homologous to 16S rRNA	0.18
976	AL112360	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen deprivation	0.22
		Danaus chrysippus orientis 12S ribosomal RNA gene, partial	
977	AF389904	sequence; mitochondrial gene for mitochondrial product	0.022
979	X96999	P.vulgaris Ypr10 gene	1.5
980	AL110480	Caenorhabditis elegans cosmid Y24F12A, complete sequence	0.16
981	AJ387746	Weldenia candida chloroplast tRNA-Leu intron and trnL-F intergenic spacer	1.8
		Plasmodium falciparum chromosome 2, section 14 of 73 of the	
982	AE001377	complete sequence	0.002
983	U26946	Gallus gallus inhibin beta A subunit mRNA, complete cds	2.1
984	AC091651	Homo sapiens clone RP11-17D6, complete sequence	2E-76
985	AF143382	Arabidopsis lyrata pistillata gene, complete cds	0.02
		E.histolytica plasmid genes for ribosomal RNA and hemolysins	
986	Z29969	HLY1, HLY5mc1 HLY5mc2 HLY4	0.02
987	XM_001301	Homo sapiens KIAA0469 gene product (KIAA0469), mRNA	0.022
		Bacillus subtilis genomic DNA containing gutA to cotA region, 48	
988	AB007638	degree	0.021
989	XM_012394	Homo sapiens myosin IXA (MYO9A), mRNA	7E-09
990	AF258322	Zenaida galapagoensis beta fibrinogen gene, intron 7 and partial cds	5
991	XM_048778	Homo sapiens similar to coagulation factor VIII, procoagulant component (hemophilia A) (H. sapiens) (LOC93021), mRNA	0.73
992	AB056815	Macaca fascicularis brain cDNA clone:QflA-15307, full insert sequence	6.4

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Homo sapiens partial SLC22A3 gene for organic cation transporter 3,	
993	AJ403948	exons 3-4	0.008
		Ephedrus laevicollis NADH dehydrogenase 1 gene, mitochondrial	
994	AF069188	gene encoding mitochondrial protein, partial cds	0.082
		Staphylococcus aureus quinone oxidoreductase and glycine betaine	
995	AF282920	transporter genes, complete cds	6.6
000	4 D021205	Homo sapiens gene for calcium-sensing receptor, exons, promoter	
996	AB031325	region	0.081
997	AE001421	Plasmodium falciparum chromosome 2, section 58 of 73 of the complete sequence	0.74
997	AE001421	Rattus norvegicus phosphoinositide-specific phospholipase C epsilon	0.74
998	AF323615	mRNA, complete eds	2.3
226	AF323013	Inicia, complete cus	2.3
999	AF305103	Picea abies clone 1PABC5D3 hypermethylated genomic sequence	0.25
	111303103	Oryza sativa microsatellite MRG4442 containing (TC)X17, closest to	0.20
1000	AY022117	marker RG450, genomic sequence	0.7
1001	NM 023324	Mus musculus pellino 1 (Peli1), mRNA	0.26
1002	X06329	Mouse repetitive sequence L1Md-F type 5 'end (1G-6)	0.73
1003	AE001173	Borrelia burgdorferi (section 59 of 70) of the complete genome	0.24
1004	AK021535	Homo sapiens cDNA FLJ11473 fis, clone HEMBA1001712	0.25
1005	U90529	Plasmodium vivax SV-5 Pv200 high-binding region I gene, partial cds	0.24
		Homo sapiens MAD2 (mitotic arrest deficient, yeast, homolog)-like 1	
1006	XM_003560	(MAD2L1), mRNA	4E-26
		Homo sapiens isolate HERV-H-PA9 long terminal repeat, partial	
1007	AF315106	sequence	0.000001
1008	U25177	Helobdella robusta cyclin A mRNA, partial cds	0.025
1009	NM_021401 AF339813	Mus musculus secreted and transmembrane 1 (Sectm1), mRNA	0.084
1010	AF339813	Homo sapiens clone IMAGE:297403, mRNA sequence	- Z
1011	AF020599	Equus caballus insulin-like growth factor 2 (IGF2) gene, partial cds	0.38
1012	BC001815	Homo sapiens, clone IMAGE:2989261, mRNA	7E-43
1012	Beddidib	Frankia sp. ArI3 cryptic plasmid pFQ31 ORF AF, korSAF gene, ORF	72.0
		CrF, ORF DrF, repF gene, ORF FrF, ORF GF, ORF HF, ORF JF,	
		ORF KrF, ORF LrF, ORF MF, ORF NrF, ORF OF, ORF PrF, parAF	ĺ
1013	AJ297945	gene and ORF SrF	0.64
1014	Z71327	S.cerevisiae chromosome XIV reading frame ORF YNL051w	0.61
		Homo sapiens hypothetical gene supported by AB007970	
1015	XM_031156	(LOC90360), mRNA	0.003
1016	XM_007274	Homo sapiens hypothetical protein FLJ10242 (FLJ10242), mRNA	0.029
		Helicobacter pylori, strain J99 section 56 of 132 of the complete	
1017	AE001495	genome	0.076
1018	AK024606	Homo sapiens cDNA: FLJ20953 fis, clone ADSE01979	e-109
	170.00.0	Melaleuca lateritia 5S ribosomal RNA gene, partial sequence; and 5S	١.,
1019	AF049434	ribosomal RNA intergenic spacer, complete sequence	1.9

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
1020	AF078543	Pestivirus type 4 strain MN fetus polymerase (NS5b) gene, partial cds	0.68
1021	M63636	Streptococcus thermophilus beta-D-galactosidase (lacZ) gene, complete cds	0.6
1022	AJ270226	Entodinium caudatum partial mRNA for putative glycosyltransferase, clone L48	0.67
1023	M59429	T. thermophila F-antigen (tfa) gene, complete cds	0.009
1024	X77801	S.beecheyi genomic DNA with direct repeats and integrated Hepatitis B virus	1.9
1025	AL590109	Human DNA sequence from clone RP11-405J10 on chromosome 10, complete sequence [Homo sapiens]	6E-84
1026	AF269904	Staphylococcus epidermidis strain SR1 clone step.1029h10 genomic sequence	0.076
1027	AL591343	Human DNA sequence from clone RP11-260H5 on chromosome 6, complete sequence [Homo sapiens]	0.22
1028	Z12986	O.aries rearranged T-cell receptor delta	0.78
1029	AY042836	Arabidopsis thaliana Unknown protein mRNA, complete cds	0.25
1030	X00117	Galago Alu repeat type I, GAL9	0.0001
1031	XM_015858	Homo sapiens Werner syndrome (WRN), mRNA	6.6
1032	M27444	Bos taurus (clone pTKD7) dopamine and cyclic AMP-regulated neuronal phosphoprotein (DARPP-32) mRNA, complete cds	0.63
1033	AF272852	Arabidopsis thaliana beta 1,2-xylosyltransferase mRNA, complete cds	0.77
1034	L08174	Romanomermis culicivorax mitochondrial NADH dehydrogenase subunits 6 and 3 (ND6, ND3) cytochrome P450-like protein genes, complete CDS	0.084
1035	AF387598	Nicotiana tabacum EPSP synthase 1 (EPSPS1) gene, partial cds	0.007
1036	AF157094	Sarcophilus harrisii mitochondrial DNA control region, partial sequence	0,23
1037	XM_046761	Homo sapiens hypothetical gene supported by AK000735 (LOC92705), mRNA	0.66
1038	AF348479	Bos taurus myostatin gene, partial cds	6.1
1039	XM_050269	Homo sapiens solute carrier family 1 (glial high affinity glutamate transporter), member 3 (SLC1A3), mRNA	0.21
1040	AF164967	Canine distemper virus strain A75/17, complete genome	7.1
1041	AB028634	Flammulina velutipes mitochondrial gene for RNA polymerase, DNA polymerase, complete and partial cds	0.086
1042	AF312925	Homo sapiens cryptic (CFC1) gene, complete cds	0.000004
1043	U67596	Methanococcus jannaschii section 138 of 150 of the complete genome	0.26
1044	L04631	Human cholesterol 7 alpha-hyroxylase (CYP7) gene, exon 3	0.009
1045	U50336	Mus musculus Ah receptor gene, promoter region and exon 1, partial sequence	2.1
1046	AF105001	Mus musculus Fab1098 immunoglobulin light chain variable region mRNA, partial cds	0.8

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
1047	U55276	Rattus norvegicus microtubule-associated protein 1B gene, promoter region	0.79
1048	AF056302	Drosophila melanogaster eIF-2alpha kinase (GCN2) mRNA, complete cds	0,63
1049	NC_001322	Drosophila yakuba mitochondrion, complete genome	0.24
		Oreina coerulea cytochrome oxidase subunit I gene, partial cds;	
1050	AF097036	mitochondrial gene for mitochondrial product	0.23
		Pasteurella haemolytica-like sp. pllktC, pllktB, pllktD, and leukotoxin	
1051	L12148	structural protein (pllktA) genes, complete cds	0.21
		Mus musculus DNA for DNA dependent protein kinase catalytic	
1052	AB005153	subunit, exon 5, 6, 7	0.006
		Brassica oleracca var. italica floral homeotic protein pseudogene,	
1053	AF126733	partial sequence	2.1
1054	AF090900	Homo sapiens clone HQ0189 PRO0189 mRNA, complete cds	0.000001
1055	AJ223260	Chromodoris purpurea mitochondrial COI gene, partial	0.007
		Pseudoleistes virescens cytochrome b (cytb) gene, mitochondrial gene	
1056	AF089052	encoding mitochondrial protein, partial cds	0.77
1057	AL138748	Human DNA sequence from clone RP4-732E19 on chromosome Xq21.1-21.33 Contains GSSs, complete sequence [Homo sapiens]	0.59
1058	AL132767	Human DNA sequence from clone RP5-819L10 on chromosome 6q12- 13. Contains GSSs, complete sequence [Homo sapiens]	5
1059	AF032670	Lucilia cuprina Notch homolog (Scl) gene, exon B	0.055
		Plasmodium falciparum chromosome 2, section 8 of 73 of the	
1060	AE001371	complete sequence	0.062
1061	AY039845	Arabidopsis thaliana At1g35160/T32G9_30 mRNA, complete cds	4.6
		Sindoropsis le-testui tRNA-Leu (trnL) gene, partial intron sequence;	
1062	AF365189	chloroplast gene for chloroplast product	0.11
1063	U60591	Drosophila melanogaster kuzbanian (kuz) mRNA, complete cds	0.41
1064		Homo sapiens SDS-stable vimentin-bound DNA fragment	
1064	AJ297052	HEF42VIM12	4E-09
1065	AB023191	Homo sapiens mRNA for KIAA0974 protein, partial cds	0.23
1066	AK021989	Homo sapiens cDNA FLJ11927 fis, clone HEMBB1000402	0.33
1067	XM_038524	Homo sapiens hypothetical protein FLJ23059 (FLJ23059), mRNA	e-169
1068	XM_045967	Homo sapiens estrogen receptor 1 (ESR1), mRNA	4.9
1069	AF354258	Homo sapiens polynucleotide kinase-3'-phosphatase (PNKP) gene, complete cds	0.66
1070	AJ404320	Posidonia oceanica medium repetitive sequence, clone pPopm6	0.038
1071	NC 002756	Mertensiella luschani mitochondrion, complete genome	4.9
		Human DNA sequence from cosmid L21F12, Huntington's Disease	
1072	Z68886	Region, chromosome 4p16.3	2E-10
		Plasmodium falciparum chromosome 2, section 7 of 73 of the	
1073	AE001370	complete sequence	0.025

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
1074	XM 033521	Homo sapiens hypothetical protein FLJ10357 (FLJ10357), mRNA	
1075	U02239	Mycoplasma genitalium random genomic clone xd5, partial cds	1.5
1073	002239		0.006
1076	AE006395	Lactococcus lactis subsp. lactis IL1403 section 157 of 218 of the complete genome	0.32
		Oryza sativa microsatellitc MRG6151 containing (ATAC)X8,	
1077	AY023826	genomic sequence	0.031
1079	X63213	B.taurus CI-B12 mRNA for ubiquinone oxidoreductase complex	7
1080	AK021648	Homo sapiens cDNA FLJ11586 fis, clone HEMBA1003720	3E-13
		Dictyostelium discoideum ABC transporter mdrA1 (MDRA1) and	
1081	AF246689	ABC transporter mdrA2 (MDRA2) genes, complete cds	0.67
1082	AB016100	Oryza sativa LINE retrotransposon, endonuclease region of RILN5	0.068
		Orestias species mitochondrial control region, allele SRT1, partial	
1083	U93157	sequence	6.9
		Caenorhabditis elegans transmembrane protein NDG-4 (ndg-4)	
1084	AF173373	mRNA, complete cds	1.8
		Turkey coronavirus genomic RNA for partial spike glycoprotein (s	
		gene), 3a protein (gene 3), 3b protein (gene 3), E protein (gene 3) and	
1086	AJ310640	partial M protein (gene 4), strain turkey/UK/412/00	2
1087	M21953	Duck hepatitis B virus, complete genome	0.086
		Homo sapiens stratum corneum chymotryptic enzyme (KLK7) gene,	
1088	AF332583	complete cds, alternative transcripts	2.1
1089	AY037894	Glomus intraradices phosphate transporter mRNA, partial cds	0.084
		Rattus norvegicus a disintegrin and metalloproteinase domain 1	
1091	NM_020078	(fertilin alpha) (Adam1), mRNA	0.51
1092	AF282531	Homo sapiens clone 18qtel_c30t7 sequence	0.003
		Ichthyophthirius multifiliis immobilization antigen isoform (IAG52A)	
1093	AF324424	gene, complete cds	0.56
1094	AJ404320	Posidonia oceanica medium repetitive sequence, clone pPopm6	0.075
1096	Z46803	V.narbonensis mRNA for legumin A precursor	2.2
		Homo sapiens similar to hypothetical protein FLJ14751 (H. sapiens)	
1097	XM_050195	(LOC93274), mRNA	5.7
1098	AJ246004	Homo sapiens RET gene, 5' genomic fragment	0.00004
		HIV-1 isolate NC4057-1999 from USA pol polyprotein (pol) gene,	
1099	AY031051	partial cds	0.23
1100	AE002146	Ureaplasma urealyticum section 47 of 59 of the complete genome	0,26
		H. sapiens (D1S436) DNA segment containing (CA) repeat, clone	
1101	Z23650	AFM217zc3; single read	0.001
1102	D73382	Sugarbeet mitochondrial gene for ORFB, complete cds	0.73
1103	U67503	Methanococcus jannaschii section 45 of 150 of the complete genome	0.69
1103	007303	Human DNA sequence from clone RP11-22P2 on chromosome Xq23-	0.09
1104	AL590376	24, complete sequence [Homo sapiens]	0
1106	D14636	Mouse mRNA for PEBP2a1 protein, complete cds	0.61
	217020	process and a sol i Esti Sat process, complete our	0.01

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		The state of the global (Flash 173, Collottic)	T
NO	ACCESSN	DESCRIP	P VALUE
1107	Y14622	Mus musculus Tnfr2 gene, exon 4 to exon 8	0.71
1108	AF130049	Homo sapiens clone FLB3411 PRO0852 mRNA, complete cds	0.065
1109	AJ011009	Glycine max cytosolic glutamine synthetase (gs15) gene promoter	0.007
1110	XM_043523	Homo sapiens Rho guanine exchange factor (GEF) 11 (ARHGEF11), mRNA	0.7
1111	Z77974	H. sapiens flow-sorted chromosome 6 HindIII fragment, SC6pA5D12	0.001
1112	AJ002513	Schizosaccharomyces pombe cmb1 gene	8.2
1113	AF098878	Homo sapiens APAF1 gene, exons 6 and 7	1.9
1114	M14080	Herpesvirus saimiri thymidylate synthase gene, complete eds	0.023
1115	U44838	Glycine max extensin (SbHRGP3) gene, complete cds	0.76
1116	Y15491	Pongo pygmaeus gene encoding fertilin alpha	2.1
1117	U43145	Plasmodium chabaudi repeat organellar protein gene, complete eds	0.25
1118	Z81524	Caenorhabditis elegans cosmid F32H5, complete sequence	0.009
1119	AF052957	Homo sapiens type XV collagen (COL15A1) gene, exons 3 and 4	2.2
1120	U67523	Methanococcus jannaschii section 65 of 150 of the complete genome	0.001
1121	AL121807	S.pombe chromosome III cosmid c132	0.19
1122	X59280	S. douglasii gene for cytochrome b	0.29
1123	XM_030915	Homo sapiens KIAA0673 protein (KIAA0673), mRNA	5E-19
1124	X99254	P.falciparum gene encoding primase, small subunit	0.084
1125	AC004214	Homo sapiens clone UWGC:m32agap from 6p21, complete sequence	0.009
		Agrocybe aegerita B type DNA polymerase (Mtpol) gene, complete cds; tRNA-Asn gene, complete sequence; and unknown genes,	
1126	AF061244	mitochondrial genes for mitochondrial products	0.001
1127	AK001125	Homo sapiens cDNA FLJ10263 fis, clone HEMBB1000991	0
1128	L41046	Pisum sativum endo-1,4-beta-glucanase (EGL1) gene, complete eds	0.0001
1129	XM_008099	Homo sapiens integrin, alpha L (antigen CD11A (p180), lymphocyte function-associated antigen 1; alpha polypeptide) (ITGAL), mRNA	2E-18
1130	AE000049	Mycoplasma pneumoniae M129 section 31 of 63 of the complete genome	0.75
1131	AY019627	Oryza sativa microsatellite MRG1952 containing (AT)X40, closest to marker L128, genomic sequence	0.024
1132	Z46383	Bovine herpesvirus type 4 DNA for nonconserved region A	4.3
1133	Y07893	S.cerevisiae DNA fragment, right arm of chromosome VII	4.2
1134	XM_040867	Homo sapiens sperm acrosome associated 1 (SPACA1), mRNA	0.003
1135	XM_038693	Homo sapiens solute carrier family 9 (sodium/hydrogen exchanger), isoform 2 (SLC9A2), mRNA	2
1136	NM_033074	Mus musculus DNA segment, Chr 15, Wayne State University 59, expressed (D15Wsu59e), mRNA	0.64

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
1137	M97168	Homo sapiens X (inactive)-specific transcript (XIST) complete exon	3,4
1138	D13973	Dictyostelium discoideum DNA for Dp87 protein, complete cds	0,075
1139	AF303254	Caenorhabditis elegans gene XL153 mRNA, complete cds	2
1140	AB009926	Cyphonoccrus ruficollis mitochondrial DNA for 16S rRNA, partial sequence	0.24
1141	NC_002660	Cochliomyia hominivorax mitochondrion, complete genome	0.0009
1142	AL096756	Human DNA sequence from clone RP3-477J10 on chromosome 22q13.31-13.33 Contains a GSS, complete sequence [Homo sapiens]	0.26
1143	XM_031937	Homo sapiens hypothetical protein FLJ11183 (FLJ11183), mRNA	2.2
1144	AE003025	Drosophila melanogaster genomic scaffold 142000013385389, complete sequence	0.22
1145	Z11702	V.faba mRNA for polyphenol oxidase	0.76
1146		Homo sapiens calcium binding protein 5 (CABP5), mRNA	5,6
1147	XM 047707	Homo sapiens KIAA1265 protein (KIAA1265), mRNA	1.6
1148	AY005256	Xylocopa erythrina cytochrome b (cytb) gene, partial cds; mitochondrial gene for mitochondrial product	0.086
1151	Z96679	H.sapiens telomeric DNA sequence, clone 7PTEL026, read 7PTEL0026.seq	9E-34
1152	AJ001677	Branchiostoma floridae mRNA for phenylalanine hydroxylase, partial	0.067
1153	J03252	Human alkaline phosphatase (ALPP) gene, complete cds	0.65
1154	AK007179	Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:1700112J16, full insert sequence	0.24
1155	AK004093	Mus musculus 18 days embryo cDNA, RIKEN full-length enriched library, clone:1110034K09, full insert sequence	6.7
1156	AF370142	Arabidopsis thaliana unknown protein (T5I7.1/At2g39710) mRNA, complete cds	1.7
1157	AF053649	Homo sapiens cellular apoptosis susceptibility protein (CSE1) gene, exons 15 and 16	0.006
1158	AL080199	Homo sapiens mRNA; cDNA DKFZp434E082 (from clone DKFZp434E082)	0.081
1159	AF267225	Candidatus Carsonella ruddii natural-host Tainarys sordida ATP synthase alpha subunit (alpA) gene, partial cds; ATP synthase gamma subunit (alpG) gene, complete cds, and ATP synthase beta subunit (alpD) gene, partial cds	0.15
1160	AE000654	Helicobacter pylori 26695 section 132 of 134 of the complete genome	0.51
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
1161	AK007179	library, clone:1700112J16, full insert sequence	0.25
1162	NM_008787	Mus musculus pericentrin (Pcnt), mRNA	1.7
1163	AK024592	Homo sapiens cDNA: FLJ20939 fis, clone ADSE01583	0.077
1164	D13197	Mouse gene for immunoglobulin diversity region D1	0.076

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID	··	Tourset Toughout (Daniel 176) Genouncy	T
NO	ACCESSN	DESCRIP	P VALUE
		Plasmodium falciparum chromosome 2, section 18 of 73 of the	- 11200
1165	AE001381	complete sequence	0.073
1166	X68360	M.fascicularis gene for apolipoprotein A-II	0,0009
1167	X73032	B. juncea ribosomal intergenic spacer	0.023
		Homo sapiens hypothetical gene supported by AL137266	
1168	XM_037745	(LOC91332), mRNA	0.21
1169	NM 017755	Homo sapiens hypothetical protein FLJ20303 (FLJ20303), mRNA	0.58
1170		Homo sapiens cofilin 2 (muscle) (CFL2), mRNA	2
	_	Lactococcus lactis subsp. lactis IL1403 section 111 of 218 of the	
1171	AE006349	complete genome	0.12
1172	VA 020627	Home coniene C matein counted assessed binary 7 (CDDV7) PNA	16
11/2	AM_030037	Homo sapiens G protein-coupled receptor kinase 7 (GPRK7), mRNA Mus musculus 6 days neonate head cDNA, RIKEN full-length	1.6
1173	AK017298	enriched library, clone:5430411K23, full insert sequence	0.18
11/3	ALK017298	Homo sapiens transcription factor HOXD12 (HOXD12) and	0.18
1174	AF154915	transcription factor HOXD11 (HOXD11) genes, complete cds	0.52
11/4	711154715	Lepidopilum scabrisetum ribosomal protein system 4 (rps4) gene.	0.32
1175	AF143066	partial cds	0.51
		Human DNA sequence from clone RP11-49A4 on chromosome 6,	
1176	AL391814	complete sequence [Homo sapiens]	0.06
1177	AF003511	Toxoneuron sp. 16S ribosomal RNA gene, partial sequence	0.0000001
1178	AK022350	Homo sapiens cDNA FLJ12288 fis, clone MAMMA1001783	0,00002
		Mus musculus small inducible cytokine A21b (Scya21b) gene,	
1180	AF307986	complete cds	0.021
1181	NM_022242	Rattus norvegicus niban protein (Niban), mRNA	5
		H.sapiens CpG island DNA genomic Msel fragment, clone 173c2,	
1182	Z57380	forward read cpg173c2.ft1a	0.13
1183	XM 043937	Homo sapiens hypothetical protein MGC13033 (MGC13033), mRNA	6.4
1184	L15439	Geukensia demissa neuropeptide mRNA, complete cds	0.24
		Proechimys cayennensis mitochondrial cytb gene for cytochrome b	
1185	AJ251399	(specimen voucher V-815)	0.7
1186	U00053	Caenorhabditis elegans cosmid K03F8	0.62
	-		
		Human (genomic clones lambda-[SK2-T2, HS578T]; cDNA clones	
1187	J00277	RS-[3,4, 6]) c-Ha-ras1 proto-oncogene, complete coding sequence	7E-55
1189	X70677	X.laevis XFG 5-1 mRNA for zinc finger protein	0.23
		Saccharomyces cerevisiae mitochondrial tRNA-Tyr, tRNA-Asn, &	
1190	AJ223323	tRNA-Met genes	0.083
1191	XM_030479	Homo sapiens KIAA1462 protein (KIAA1462), mRNA	0.76
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
1192	AK016425	library, clone:4931403E03, full insert sequence	0.00003
1193	AF092854	Saguinus mystax visual pigment gene, P562 allele, intron 4	0.0001
1194	AC025732	Homo sapiens BAC clone RP11-116J19 from Y, complete sequence	1.7

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
1195	AE002177	Chlamydophila pneumoniae AR39, section 13 of 94 of the complete genome	0.011
1196	M94732	Plasmodium falciparum 15 kDa vesicular-like antigen gene, exons 1 through 4	0.009
1197	X76311	H. sapiens endothelial nitric oxidase synthase gene, exons 15 and 16	0.6
1198	L81756	Homo sapiens (subclone 1_g5 from P1 H16) DNA sequence, complete sequence	0.061
1199	Z82625	R.prowazekii genomic DNA fragment (clone A396F)	0.76
1200	XM_012528	Homo sapiens protein kinase C, beta 1 (PRKCB1), mRNA	2.1
1201	M27082	S.cerevisiae GCN2 gene (which couples GCN4 expression to amino acid availability), complete cds	0.21
1202	AK007856	Mus musculus 10 day old male pancreas cDNA, RIKEN full-length enriched library, clone:1810054D07, full insert sequence	6E-73
1203	M69188	Cotton legumin A D-genome alloallele gene, complete cds	0.076
1204	AB043132	Thunnus thynnus DNA, microsatellite Ttho-6*, partial sequence	0.001
1205	AF019084	Homo sapiens keratin 2e (KRT2E) gene, complete cds	6
1206	U32784	Haemophilus influenzae Rd section 99 of 163 of the complete genome	0,026
1207	XM_004699	Homo sapiens ubiquitin-conjugating enzyme E2H (homologous to yeast UBC8) (UBE2H), mRNA	2
1208	AF375643	Pongo pygmaeus HERV-K-GC1 preintegration site sequence	2.2
1209	AK007179	Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:1700112J16, full insert sequence	0,22
1210	AF182214	Glycine max glutamine synthetase gamma 2 subunit gene, 5'-flanking region and partial cds	0.23
1211	AF170224	Alloplectus meridensis ribulose 1,5-bisphosphate carboxylase large subunit gene, partial cds; chloroplast gene for chloroplast product	6.5
1212	NM_022978	Homo sapiens small EDRK-rich factor 1B (centromeric) (SERF1B), mRNA	0.23
1213	U17787	Gromphadorhina portentosa mitochondrion 16S ribosomal RNA, partial sequence	0.027
1214	AF144086	Papio hamadryas anubis chromosome 13 repeat region	0.25
1215	XM 010106	Homo sapiens DKFZP586N0819 protein (DKFZP586N0819), mRNA	0.73
1216	AF198964	Arabidopsis thaliana ATP sulfurylase gene, complete cds	0.072
		Xenopus laevis mRNA for winged helix transcription factor (foxd3b	0,072
1217	AJ298866	gene)	0.2
		Uroleucon obscurum large subunit ribosomal RNA gene, partial sequence; tRNA-Val gene, complete sequence; and small subunit	
1218	AF069111	ribosomal RNA gene, partial sequence	0.003
1219	X16715	D. melanogaster gene for chorion protein s16	0.23
1220	AF245117	Mus musculus leukocyte cell-surface molecule (Ly9) gene, exon 1	0.024

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
1221	AP001310	Arabidopsis thaliana genomic DNA, chromosome 3, P1 clone:MUO22	0.23
i l		Rivulus gransabanae tRNA-Ile, tRNA-Gln, and tRNA-Met genes,	
]		complete sequence; NADH dehydrogenase subunit II gene, complete	
		cds; tRNA-Trp, tRNA-Ala, tRNA-Asn, tRNA-Cys, and tRNA-Tyr	
1,,,,,	1 Toogga	genes, complete sequence; and cytochrome c oxidase subunit I gene,	
1222	AF092375	par>	0.26
1223	U84864	HIV-1 strain M14 from USA, envelope glycoprotein (env) gene, partial cds	0.76
1223	004004	Homo sapiens similar to estrogen-related receptor alpha; estrogen	0.76
1224	XM 051684	receptor-like 1 (H. sapiens) (LOC93491), mRNA	0.57
1225	AF282391	Cucumis sativus clone B99 mitochondrial genomic sequence	2.2
1226	AK026459	Homo sapiens cDNA: FLJ22806 fis, clone KAIA2845	0.024
1220	F11020435	Spodoptera littoralis nuclear polyhedrosis virus fgf, egt genes and 3	0.024
1227	AJ003131	ORFs	0.022
1228	XM 028308	Homo sapiens helicase-moi (KIAA0928), mRNA	1.8
1000	1111_020500	Fritillaria camtschatcensis clone cam36 retrotransposon TfcI, partial	1.0
1229	AF219168	sequence	2
		Homo sapiens ataxia-telangiectasia and Rad3 related protein (ATR)	
1230	AF325699	gene, partial cds, alternatively spliced	0.069
		, , , , , , , , , , , , , , , , , , ,	
1231	AE007433	Streptococcus pneumoniae section 116 of 194 of the complete genome	7.9
1232	XM_007429	Homo sapiens CGI-35 protein (LOC51077), mRNA	1E-08
		Dianous nitidulus cytochrome b (cytb) gene, mitochondrial gene	
1233	AF021084	encoding mitochondrial protein, partial cds	0.07
		Plasmodium falciparum DNA *** SEQUENCING IN PROGRESS	
1234	Z97349	*** from contig 3-06, complete sequence	2.1
1235	XM_038588	Homo sapiens like-glycosyltransferase (LARGE), mRNA	5.5
1236	X95276	P.falciparum complete gene map of plastid-like DNA (IR-B)	0,008
		Drosophila melanogaster CREB-binding protein homolog mRNA,	
1237	U88570	complete cds	1.7
1238	AK024332	Homo sapiens cDNA FLJ14270 fis, clone PLACE1004491	0.68
1239	AC006678	Caenorhabditis elegans cosmid R05G9, complete sequence	1.8
		Lycopersicon esculentum peptide deformylase-like protein mRNA,	
1240	AF250958	complete cds	0,076
1241	M26650	Human islet amyloid polypeptide (hIAPP) gene, complete cds	1.9
1242	XM_037062	Homo sapiens DKFZP564G202 protein (DKFZP564G202), mRNA	0.61
ا ا	A 170000005	Drosophila melanogaster genomic scaffold 142000013385389,	0.005
1243	AE003025 U39701	complete sequence	0.027
1244	039/01	Mycoplasma genitalium section 23 of 51 of the complete genome	2.1
1245	AE000036	Mycoplasma pneumoniae M129 section 44 of 63 of the complete genome	0.1
1243	AE000030	Methanobacterium thermoautotrophicum from bases 394245 to	8.2
1246	AE000830	404874 (section 36 of 148) of the complete genome	0.67
1240	120000000	TOTO / T (SOCIOII 30 OI 140) OI IIIC COMPLETE BEHOME	0.07

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ano r		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
1247	AL358951	Human DNA sequence from clone RP3-456L16 on chromosome 6, complete sequence [Homo sapiens]	0.64
1248	XM_045757	Homo sapiens chloride channel 4 (CLCN4), mRNA	0.56
1249	U59683	Nicotiana tabacum squalene synthase (TSS) gene, complete cds	0,005
1250	AK010434	Mus musculus ES cells cDNA, RIKEN full-length enriched library, clone:2410006O21, full insert sequence	4.9
1251	XM_045757	Homo sapiens chloride channel 4 (CLCN4), mRNA	0.61
1252	AE000717	Aquifex aeolicus section 49 of 109 of the complete genome	0.024
1253	AC024780	Caenorhabditis elegans cosmid Y43B11AR, complete sequence	0.027
1254	AF343030	Human herpesvirus 7 isolate CR18 glycoprotein H gene, complete cds	0.009
1255	AK022153	Homo sapiens cDNA FLJ12091 fis, clone HEMBB1002582	0,000000
1256	AF309947	Dictyostelium discoideum Rac1A (rac1A) gene, complete cds; and unknown gene	0.009
1257	AB051519	Homo sapiens mRNA for KIAA1732 protein, partial cds	0.25
1258	AC006681	Caenorhabditis elegans clone R13H9, complete sequence	2.2
1260	Y11648	Campylobacter jejuni waaC, galE, wla[B,C,D,E,F,G,H,I,K,L,M], cheY genes and orf1	0.25
1261	AB009592	Oryza sativa gene for cytosolic glutathione reductase, complete cds	2.1
1262	X83997	C.parasitica eapC gene	0.22
1263	AJ404320	Posidonia oceanica medium repetitive sequence, clone pPopm6	0.034
1264	AF038620	Caenorhabditis elegans cosmid H10D12	0,063
1265	U05822	Human proto-oncogene BCL3 gene, exon 2	0.000000
1266	Z79343	H. sapiens flow-sorted chromosome 6 TaqI fragment, SC6pA4G2	0.13
1267	S60289	LeB4=legumin {5' region} [Vicia faba, Genomic, 1222 nt]	0,0003
1268	AE006303	Lactococcus lactis subsp. lactis IL1403 section 65 of 218 of the complete genome	1.9
1269	AE001410	Plasmodium falciparum chromosome 2, section 47 of 73 of the complete sequence	0.008
1270	X96972	D.buzzatii copia element DNA	0.2
1271	AB042973	Nicotiana tabacum wizz gene, 5' flanking region, partial cds	0.073
1272	Z95309	Caenorhabditis elegans cosmid H36L18, complete sequence	3.5
1273	AF091745	Homo sapiens exonuclease I (EXOI) gene, exon 5	0.2
1274	AF386945	Arabidopsis thaliana Rubisco subunit binding-protein beta subunit (T5A14.11) mRNA, complete cds	0.7
1275	AF269376	Staphylococcus epidermidis strain SR1 clone step.1002h01 genomic sequence	0.076
1276	AK012864	Mus musculus 10, 11 days embryo cDNA, RIKEN full-length enriched library, clone:2810036H5, full insert sequence	0.028
1277	Z74204	S.cerevisiae chromosome IV reading frame ORF YDL156w	0.18
1278	AB049851	Macaca fascicularis brain cDNA, clone:QnpA-18831	2E-91
1279	AE001761	Thermotoga maritima section 73 of 136 of the complete genome	0.19
1280	AE002102	Ureaplasma urealyticum section 3 of 59 of the complete genome	5.5
1281	AK021982	Homo sapiens cDNA FLJ11920 fis, clone HEMBB1000312	0.006

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
1282	AE007504	Streptococcus pneumoniae section 187 of 194 of the complete genome	0.21
1283	X79095	F.trinervia pdk gene for pyruvate,orthophosphate dikinase	0.22
1284	AE007413	Streptococcus pneumoniae section 96 of 194 of the complete genome	1.7
1285	AB050513	Macaca fascicularis brain cDNA, clone:QnpA-17049	0.41
1286	AF339809	Homo sapiens clone IMAGE:274150, mRNA sequence	0.52
1287	AF074991	Homo sapiens full length inscrt cDNA YH88A03	4E-09
1288	AF277581	Campostoma anomalum microsatellite CA9 sequence	1.6
1289	AF035245	Bacillus lynceorum strain pVIL31 Bag320 satellite DNA	0.23
1291	AF298208	Dictyostelium discoideum transposon thug-T, complete sequence	0.028
		Homo sapiens mRNA; cDNA DKFZp434O2017 (from clone	
1292	AL133656	DKFZp434O2017)	0.021
1293	AK024666	Homo sapiens cDNA: FLJ21013 fis, clone CAE05223	9E-34
1294	U55367	Caenorhabditis elegans cosmid C55H1, complete sequence	2.1
1295	X16715	D. melanogaster gene for chorion protein s16	0.25
1296	AC001080	Homo sapiens (subclone 3_g7 from PAC H74) DNA sequence, complete sequence	0.69
1297	XM_051200	Homo sapiens hypothetical protein MGC5149 (MGC5149), mRNA	0.58
1298	AE000016	Mycoplasma pneumoniae M129 section 1 of 63 of the complete genome	0.026
1299	AF264059	Homo sapiens RAPD fragment lost in a grade 2 astrocytoma	0.73
1300	AE002308	Chlamydia muridarum, section 39 of 85 of the complete genome	0.078
1301	U58745	Caenorhabditis elegans cosmid C10G6, complete sequence	0.26
1302	XM_007768	Homo sapiens KIAA1018 protein (KIAA1018), mRNA	2.2
1303	AC000102	Homo sapiens Chromosome 22q11.2 BAC Clone 60b5 In GNAZ-BCR Region, complete sequence	0.17
1304	AF338734	Homo sapiens hypothetical PHD zinc finger protein XAP135 pseudogene, complete sequence	2.3
1305	AF211141	Carsonella ruddii natural-host Pachypsylla celtidis tryptophanyl-tRNA synthetase (trpS) gene, partial cds; 16S ribosomal RNA, 23S ribosomal RNA, and 5S ribosomal RNA genes, complete sequence, transaldolase (tnl) and alkyl hydroperoxide reductase small >	0.087
1306	L81391	Homo sapiens (subclone 2_a6 from P1 H39) DNA sequence, complete sequence	0.18
1307	AB013613	Oryza sativa DNA, centromere sequence RCB11	0.085
1308	U03906	Macroscelides proboscideus aldehyde dehydrogenase I, eta-crystallin mRNA, complete cds	0.26
1309	L13470	Human thyroxine-binding globulin gene, complete cds	0.21
1310	AK026225	Homo sapiens cDNA: FLJ22572 fis, clone HSI02313	3E-27
1311	XM 030391	Homo sapiens dJ467N11.1 protein (DJ467N11.1), mRNA	0.001
1312	XM_017923	Homo sapiens hypothetical protein FLJ12660 (FLJ12660), mRNA	2E-32

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
1313	M57256	B.hermsii vmp21 gene encoding Vmp21 outer membrane lipoprotein	0.58
1314	Y14051	Staphylococcus aureus mecA, mecR1, mecI genes and ORF168, ORF142, ORF44, ORF145 and ORF224	0.081
1315	XM_032018	Homo sapiens similar to RIKEN cDNA 2310040G17 gene (M. musculus) (LOC90480), mRNA Octodon lunatus cytochrome b gene, partial cds; mitochondrial gene	0.027
1316	AF227514	for mitochondrial product Plasmodium falciparum chromosome 2, section 51 of 73 of the	1.9
1317	AE001414 Z21622	Complete sequence S. cerevisiae RIB7 gene encoding HTP reductase	0.73 0.77
1319	XM 040462	Homo sapiens ATP-binding cassette, sub-family D (ALD), member 3 (ABCD3), mRNA	1.8
1320	Z65229	H.sapiens CpG island DNA genomic Mse1 fragment, clone 28a8, forward read cpg28a8.ft1a	0.23
1321	XM_040095	Homo sapiens hypothetical protein (DKFZP434G0310), mRNA	2.3
1322 1323	XM_027314 AF282064	Homo sapiens hypothetical protein FLJ22351 (FLJ22351), mRNA Homo sapiens clone 2qtel_edc17iF-c17jF sequence	1E-14 3E-88
1324	AF101098	Homo sapiens collagen type XI alpha-1 (COL11A1) gene, exon 38	1.3
1325	AB050093	Neoceratodus forsteri mRNA for pituitary glycoprotein hormone alpha subunit, complete cds	0.009
1326	AL591507	Human DNA sequence from clone RP11-289C14 on chromosome 6, complete sequence [Homo sapiens]	0.026
1327	AJ400873	Homo sapiens partial GPLD1 gene for glycosylphosphatidylinositol phospholipase D, exons 15-20 Mus musculus orphan G protein-coupled receptor (Raig2) mRNA,	1.9
1328	AF378831	lous musculus orpnan G protein-coupled receptor (Raig2) mR.NA, complete cds	1.9
1329 1330	XM_030116 X54806	Homo sapiens hypothetical protein MGC3077 (MGC3077), mRNA R.norvegicus mRNA for cytokeratin type I (3' end)	0.77
1331	XM_013141	Homo sapiens ATPase, Cu++ transporting, alpha polypeptide (Menkes syndrome) (ATP7A), mRNA	0.72
1332	NM_026428	Mus musculus RIKEN cDNA 1810027P18 gene (1810027P18Rik), mRNA	0.019
1333	U85448	Actinidia deliciosa sucrose-phosphate synthase (KSPS-1) mRNA, partial cds	0,001
1334	AK013974	Mus musculus 13 days embryo head cDNA, RIKEN full-length enriched library, clone:3110002D09, full insert sequence	0.74
1335 1336	U25641 NC 002102	Tetrahymena thermophila telomerase component p80 mRNA, complete cds	0.008
1337	M14292	Lactobacillus helveticus plasmid pLH1, complete sequence Human L1Heg repetitive element from the intergenic region of the epsilon and G-gamma globin genes	0.78 5E-08

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NO ACCESSN DESCRIP P VALUE			Table 3A Nearest Neighbor (BlastN vs. Genbank)	
Petunia axillaris haplotype S15 self-incompatibility ribonuclense mRNA, complete ods 2.1	SEQ ID			
1339 AF239910 mRNA, complete ods 0.25	NO	ACCESSN		P VALUE
1340 XM_035344 Homo sapiens KIAA1688 protein (KIAA1688), mRNA 1.6	1338	AF239910		2.1
1340 XM_035344 Homo sapiens KIAA1688 protein (KIAA1688), mRNA 1.6	1339	AF116872	Candida albicans alternative oxidase (AOX2) gene, complete cds	0.25
Strongyloides stercoralis G protein alpha submit 2 (gps.2) gene, complete cds 0.76	1340	XM_035344		1.6
AF292561	1341	XM_046636	Homo sapicns hypothetical protein FLJ20080 (FLJ20080), mRNA	6E-74
1343 AF008597 gene, complete cds 0.26	1342	AF292561		0.76
1344 AF347671 Equine rhinitis A virus isolate PERV/62 P1 (P1) gene, partial cds 6.9				
1345 U89019 Hepatitis C virus polyprotein gene, complete cds 0.27				
1346 XM_043815 Homo sapiens hypothetical protein FL/21736 (FLJ21736), mRNA 2.2 1347				
1347 L39930 Podecocyne carnea fibrillin mRNA, partial cds 0.7	1345	U89019	Hepatitis C virus polyprotein gene, complete cds	0.27
1348 Z21512 K.lactis GAL80 gene encoding KIGAL80 protein Homo sapiens (subclone 2_c10 from P1 H32) DNA sequence, complete sequence Seacharomyces cerevisiae mitochondrion transfer RNA-Met (RNA-fMet) gene, transfer RNA-Pro (RNA-Pro) gene, 9S ribosomal RNA (9S RNA) gene Q. 25 Music March (180 km sapiens Christophera (180 k	1346	XM 043815	Homo sapiens hypothetical protein FLJ21736 (FLJ21736), mRNA	2.2
1348 Z21512 K.lactis GAL80 gene encoding KIGAL80 protein 0.075	1347	L39930		
Home sapiens (subclone 2_c10 from P1 H32) DNA sequence, complete sequence 2E-11	1348	Z21512		
Saccharomyces cerevisiae mitochondrion transfer RNA-fMet (tRNA-fMet) gene, transfer RNA-fMet (tRNA-fMet) gene, transfer RNA-fMet (tRNA-fMet) gene, transfer RNA-fWet) gene, cy8 ribosomal RNA (9S rRNA) gene			Homo sapiens (subclone 2 c10 from P1 H32) DNA sequence,	
Med gene, transfer RNA-Pro (RNA-Pro) gene; 9S ribosomal RNA (0.95 rRNA) gene Mus musculus 11 days pregnant adult female ovary and uterus cDNA, RIKEN full-length enriched library, clone:5033428C03, full insert sequence 6.2	1349	AC001490	complete sequence	2E-11
1350			Saccharomyces cerevisiae mitochondrion transfer RNA-fMet (tRNA-	
Mus musculus 11 days pregnant adult female ovary and uterus cDNA, RIKEN full-length enriched library, clone:5033428C03, full insert sequence 6.2			fMet) gene; transfer RNA-Pro (tRNA-Pro) gene; 9S ribosomal RNA	
RIKEN full-length enriched library, clone:5033428C03, full insert sequence 6.2	1350	L36891		0.025
Homo sapiens DNA for choline kinase like protein and muscle type carultine palmitoyltransferase I, partial and complete cds 0.069				
AB003286 Plasmodium falciparum reticulocyte binding protein 2 homolog B gene, complete cots 0.069	1351	AK017200		6.2
1353 AF312917 Plasmodium falciparum reticulocyte binding protein 2 homolog B gene, complete ods 0.081				
335	1352	AB003286		0.069
M. thermoautotrophicum trpE, trpG, trpC, trpF, trpB, trpA, trpD genes, complete cds 0.26	1353	AF312917		0.081
3354 M65060 genes, complete cds 0.26			M. thermoautotrophicum trpE, trpG, trpC, trpF, trpB, trpA, trpD	
1355 AB025642 clone:T32G24 0.34 1356 L24957 Zygosaccharomyces cidri alpha-galactosidase gene, complete cds 0.08 1357 L38437 gene, complete cds 0.27 1358 A301686 Polystoma sp. DNA-25 ITS 1, isolate DNA-25 0.27 1359 AB002805 Homo sapicas OS-9 gene, 5' upstream region and partial cds 0.0004 1361 XM_030723 Homo sapicas OS-9 gene, 5' upstream region and partial cds 0.0004 1362 NM_031881 musculus neural precursor cell expressed, developmentally down-regulated gene 4b (Nedd4b), mlN/A 3.8 1363 AC008835 sequence 0.72 1364 NC_002660 Cochliomyvia hominivorax mitochondrion, complete genome 0.074 1365 AF072S89 Python regins alpha enolase mRNA, complete cds 1.4	1354	M65060		0.26
1356 L24957 Zygosaccharomyces cidri alpha-galactosidase gene, complete cds 0.08				
Ratius norvegicus NADH ubiquinone oxidoreductase subunit (IPI3) gene, complete cds 0.27	1333	AB025642	cione: 132/G24	0.34
1357 1.38437 gene, complete cds 0.27 1358 AJ301686 Polystoma sp. DNA-25 TTS 1, isolate DNA-25 2.2 1359 AB002805 Homo sepicans CG-9 gene, 5' upstream region and partial cds 0.0004 1361 XM 030723 Homo sepicans CGI-107 protein (LOCS1012), mRNA 3.1 Mus musculus neural precursor cell expressed, developmentally down-regulated gene 4b (Ned404b), mRNA 3.8 1363 AC008855 sequence 0.72 1364 NC 002660 Cochliomyia hominivorax mitochondrion, complete genome 0.074 1365 AF072889 Pythor regims alpha enolase mRNA, complete cds 1.4	1356	L24957		0.08
1358 AJ301686 Polystoma sp. DNA-25 TTS 1, isolate DNA-25 2.2 1359 AB002805 Homo sepicas OS-9 gene, 5' upstream region and partial cds 0.0004 3361 XM_030723 Homo sepicas CGI-107 protein (LOCS1012), mRNA 3.1 Mus musculus neural precursor cell expressed, developmentally down-regulated gene 4b (Nedd4b), mRNA 3.8 Homo sepiens chromosome 5 clone CTD-2178M23, complete sequence 0.72 1364 NC_002660 Cochliomyria hominivorax mitochondrion, complete genome 0.074 1365 AF0702S89 Python regins alpha enolase mRNA, complete cds 1.4	,,,,,,	* ***		
AB002805 Homo sapiens OS-9 gene, 5' upstream region and partial cds 0.0004				
1361 XM_030723 Homo sapiens CGI-107 protein (LOC51012), mRNA 3.1				
Mus musculus neural precursor cell expressed, developmentally down- 1362 NM_031881 regulated gene 4b (Nedd4b), mtNNA 3.8 Homo sepiens chromosome 5 clone CTD-2178M23, complete sequence 0.72 1364 NC_002660 Cochliomyria hominivorax mitochondrion, complete genome 0.074 1365 AF072S89 Python regims alpha enolase mtNA, complete cds 1.4				
1362 NM_031881 regulated gene 4b (Nedd4b), mRNA 3.8	1301	AIVI_U3U/23		3,1
Homo seplens chromosome 5 clone CTD-2178M23, complete sequence 0.72	1362	NM 031881		2.8
1363 AC008855 sequence 0.72 1364 NC 002660 Cochlionyvia hominivorax mitochondrion, complete genome 0.074 1365 AF072S89 Python regims alpha enolase mRNA, complete cds 1.4	1302	141,031001		5.0
1364 NC_002660 Cochliomyia hominivorax mitochondrion, complete genome 0.074 1365 AF072589 Python regins alpha enolase mRNA, complete cds 1.4	1363	AC008855		0.72
1365 AF072589 Python regius alpha enolase mRNA, complete cds 1.4				
	1365			
	1366		Vicia faba nod-CCP1 gene, exons 1 and 2	

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
1367	XM_011052	Homo sapiens 28kD interferon responsive protein (IFRG28), mRNA	1.6
1368	AL390157	Homo sapiens mRNA; cDNA DKFZp434D179 (from clone DKFZp434D179)	0.0005
1369	AL023822	Caenorhabditis elegans cosmid Y102A5A, complete sequence	0.046
1370	U36918	Mesocricetus auratus mucin (MUC1) mRNA, complete cds	4.4
1371	AF081786	Homo sapiens lysyl hydroxylase 1 gene, promoter region and partial cds	0.18
1372	XM_035682	Homo sapiens similar to hypothetical protein FLJ22639 (H. sapiens) (LOC91047), mRNA	8E-08
1373	Z54645	H.sapiens CpG island DNA genomic Msc1 fragment, clone 14h7, reverse read cpg14h7.rt1b	0.027
1374	AB012135	Schizosaccharomyces pombe gene for catalytic subunit (C5) of proteasome, complete cds	1.9
1375	L47352	Brassica napus (clone Sta 39-4) arabinogalactan protein mRNA, complete cds	0.73
1376	AB036993	Homo sapiens RHAG gene for Rh50 glycoprotein, partial cds	0.0000001
1377	U67535	Methanococcus jannaschii section 77 of 150 of the complete genome	0.03
1378	AF203637	Myotis myotis microsatellite A2 sequence	0.017
1379	XM_004748	Homo sapiens hypothetical protein FLJ11110 (FLJ11110), mRNA	1.1
1380	AJ249744	Campylobacter jejuni cjaE gene for putative membrane protein CjaE	0.69
1381	AL591033	Human DNA sequence from clone RP11-94L3 on chromosome 6, complete sequence [Homo sapiens]	1.7
1382	AJ242625	Mus musculus Dmp-1 gene, exons 1-6	0.71
1383	AB036851	Ciona intestinalis mRNA for pellino, complete cds	0.23
1384	XM_012972		2E-09
1385	AF175313	Oncorhynchus mykiss cardiac sodium-calcium exchanger (NCX) mRNA, complete cds	0.47
1386	AF283669	Homo sapiens kallikrein 14 (KLK14) gene, complete cds	0.44
1387	AY018560	Oryza sativa microsatellite MRG0885 containing (AT)X18, genomic sequence	0.63
1388	AK027272	Homo sapiens cDNA FLJ14366 fis, clone HEMBA1001020	0.019
		Homo sapiens roundabout (axon guidance receptor, Drosophila)	
1389	XM_003052	homolog I (ROBO1), mRNA	1E-49
1390	AF208532	Homo sapiens fatty acid omega-hydroxylase CYP4A11 (CYP4A11) gene, complete cds	0.69
1391	AF233326	Mus musculus tumor necrosis factor receptor-associated factor (Traf2) gene, exons 2, 3, and 4	0.72
1392	AE001413	Plasmodium falciparum chromosome 2, section 50 of 73 of the complete sequence	0,15
1394	X54889	Sea urchin developmentally regulated ectodermal gene	0.023

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
1395	AF378868	Arabidopsis thaliana AT5g11200/F2I11_90 mRNA, complete cds	2.5
1396	AF328539	Homo sapiens clone 12qtel_c64t3 sequence	0.0003
1397	AE006450	Lactococcus lactis subsp. lactis IL1403 section 212 of 218 of the complete genome	0.47
1398	Z12840	O.cuniculus mRNA for protein of unknown function	2E-97
1399	U31165	Rattus norvegicus SH3 domain binding protein (CR16) gene, exon 6	1.9
1400	Y15005	Bison bonasus mitochondrial cytB gene	1.8
1401	AC091267	Caenorhabditis elegans cosmid Y37B11A, complete sequence	3.7
1402	AB055364	Macaca fascicularis brain cDNA, clone:QflA-12522	e-119
1403	NC_001610	Didelphis virginiana mitochondrion, complete genome	0.006
1404	AF362372	Dictyostelium discoideum histidine kinase DhkJ (dhkJ) gene, complete cds	0.055
		Homo sapiens type II integral membrane protein (NKG2-E) gene, partial cds; and type II integral membrane protein (NKG2-F) gene,	
1405	AF027164	complete cds	3.6
1406	M76376	Human cysteine-rich protein (CRP) gene, exon 2	0.027
1407	D31863	Mouse Pig-a gene for GPI-anchor biosynthesis (PIG-A protein), exon 6 and complete cds	0.026
1408	XM_031315	Homo sapiens hypothetical protein FLJ14033 similar to hypoxia inducible factor 3, alpha subunit (FLJ14033), mRNA	0.000007
1409	AY019369	Oryza sativa microsatellite MRG1694 containing (AT)X32, genomic sequence	0.0001
1410	AF067610	Caenorhabditis elegans cosmid F41A4	0.084
1411	AJ401391	Drosophila melanogaster mRNA for PDGF/VEGF-like protein (CG7103 gene)	0.18
1412	XM_008417	Homo sapiens hypothetical protein FLJ20694 (FLJ20694), mRNA	0.0000001
1413	AF242195	Homo sapiens KLK15 (KLK15) gene, complete cds, alternatively spliced	e-160
1414	Y14077	Bacillus subitlis 10.6 Kb chromosomal DNA: glyB-prsA region	0.008
1415	AE006802	Sulfolobus solfataricus section 161 of 272 of the complete genome	0.057
1416	AY007149	Homo sapiens clone CDABP0086 mRNA sequence	0.092
		Oxytricha nova (hypotrichous ciliate) (clones LMiC2-(5,6,8))	
1417	K02628	micronuclear DNA, C2 gene (version 3), complete cds	0.063
1419	X14260	Xenopus tropicalis alpha-globin gene	1.7
1420	AJ401391	Drosophila melanogaster mRNA for PDGF/VEGF-like protein	0,15
1420	XM 032580	(CG7103 gene) Homo sapiens nebulette (NEBL), mRNA	1.3
1421	AIVI_032380		1.3
1422	XM_008060	Homo sapiens solute carrier family 7 (cationic amino acid transporter, y+ system), member 5 (SLC7A5), mRNA	2E-12
1424	AF035820	Hordeum vulgare gibberellin action negative regulator SPY mRNA, complete cds	2.1

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
1425	4 Y 0 50 5 50	Human DNA sequence from clone RP1-241K1 on chromosome 6,	
	AL353672	complete sequence [Homo sapiens]	3.8
1426	AF070565	Homo sapiens clone 24425 mRNA sequence	0.2
1427	AK026914	Homo sapiens cDNA: FLJ23261 fis, clone COL05862	0.009
1428	X14735	Podospora anserina mtDNA for large subunit rRNA	0,21
1429	XM 004081	Homo sapiens aldehyde dehydrogenase 5 family, member A1	
1430	Y10687	(succinate-semialdehyde dehydrogenase) (ALDH5A1), mRNA Sulfolobus acidocaldarius gene encoding purine NTPase	1,4
1430	11000/	Suitoloous acidocaldarius gene encoding purine N1Pase	0.019
1431	AF168678	Drosophila melanogaster adenylyl cyclase 78C-s mRNA, complete cds	
1432	AC006666	Caenorhabditis elegans cosmid H31G24, complete sequence	0.13
1433	X02473	Calf mRNA for delta subunit of muscle acetylcholine receptor	0.014
1433	202473	Staphylococcus epidermidis strain SR1 clone step.1051c08 genomic	1,1
1434	AF270148	sequence	0.006
1434	AF2/0146	scquence	0.006
1435	XM 008084	Homo sapiens hypothetical protein LOC55565 (LOC55565), mRNA	3E-36
1436	AF007876	Homo sapiens Na,K-ATPase beta 2 subunit gene, complete cds	0.008
1,50	12 001010	Mus musculus adult male testis cDNA, RIKEN full-length enriched	0.008
1437	AK016592	library, clone:4933400F21, full insert sequence	0.61
		Homo sapiens genomic DNA, chromosome 6p21.3, HLA class I	0.01
1438	AB023053	region, clone:53L9, complete sequence	2E-10
			22 10
1439	XM_035837	Homo sapiens cAMP responsive element modulator (CREM), mRNA	1.7
1440	Y12710	A.thaliana gene encoding shaggy-like kinase gamma	0.068
		Homo sapiens lipocalin-1 interacting membrane receptor (LIMR)	
1441	AF351620	gene, complete cds	0.028
1442	U67560	Methanococcus jannaschii section 102 of 150 of the complete genome	2.1
		Homo sapiens potassium voltage-gated channel, delayed-rectifier,	
1443	NM_002251	subfamily S, member 1 (KCNS1), mRNA	1.9
		Homo sapiens similar to retinoblastoma inhibiting gene 1 (M.	
1444	XM_027341	musculus) (LOC89932), mRNA	8000,0
		Arabidopsis thaliana ecotype RLD chromomethylase (CMT1) gene,	
1446	AF039373	complete cds	1.9
		Rattus norvegicus plasma membrane Ca2+ ATPase isoform 3	
1447	U29397	(PMCA3) gene, 5' flanking region	0.008
ا ا	****		
1448 1449	XM_036536 AK000677	Homo sapiens hypothetical protein FLJ10402 (FLJ10402), mRNA	4.1
1449	AK0006// AJ243538	Homo sapiens cDNA FLJ20670 fis, clone KAIA4743	1E-09
1430	AJ243538	Glomus mosscae fox2 gene for Fox2 protein	0.018
1451	AJ296103	Staphylococcus aureus repNVH99 gene for replication protein and	0.00
1431	13270103	smr gene	0.23
1452	AF084911	Bombus ternarius cytochrome b gene, partial cds; mitochondrial gene for mitochondrial product	1.4
1432	71.004311	Homo sapiens (subclone 3 g7 from PAC H74) DNA sequence.	1.4
1453	AC001080	complete sequence	0.72
1400		compacts sequence	0.72

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Chlamydomonas reinhardtii partial dhc10 gene for 1 beta dynein	
1483	AJ242525	heavy chain, exons 16-53	5.8
		Daltonia sp. mitochondrial NADH dehydrogenase, and cytochrome b	
1484	L19842	genes, 3' end, and transfer RNA-Ser gene	0.001
1485	AJ239534	Camis familiaris tRNA-derived SINE element, clone D254UC12	0.009
1486	AF044206	Homo sapiens cyclooxygenase (COX-2) gene, promoter and exon 1	0.2
		Chaitoregma tattakana mitochondrial genes for small subunit rRNA,	
1487	AB035880	tRNA-Val, large subunit rRNA, partial and complete sequences	0.23
		Pasteurella multocida PM70 section 172 of 204 of the complete	
1488	AE006205	genome	0.26
1400	70.5.011446	The section of the se	
1489	XM_011446	Homo sapiens hypothetical protein FLJ10775 (FLJ10775), mRNA	6E-16
1490	NM_030714	Mus musculus deltex 3 homolog (Drosophila) (Dtx3), mRNA	0.39
1491	AE001407	Plasmodium falciparum chromosome 2, section 44 of 73 of the complete sequence	
1491	BC005530	Mus musculus, clone IMAGE:3495762, mRNA	0.001
1492	DC003330	IVIUS IIIUSCIIIUS, CIONE IIVIAGE: 3493 /62, IIIRIVA	6.4
1493	XM 017183	Homo sapiens hypothetical protein FLJ22474 (FLJ22474), mRNA	0.0007
1494	AF198964	Arabidopsis thaliana ATP sulfurylase gene, complete cds	0.007
1495	AK026781	Homo sapiens cDNA: FLJ23128 fis, clone LNG08384	0.0002
1496	AF307860	Homo sapiens CYR61 protein (CYR61) gene, complete cds	3.9
1470	AL-307800	Human DNA sequence from clone RP11-389N9 on chromosome 6.	3.9
1497	AL591667	complete sequence [Homo sapiens]	0.004
1498	XM 002800	Homo sapiens kinesin-like protein 2 (hklp2), mRNA	2E-53
1499	M90970	Atriplex numnularia DNA sequence	4.8
1477	14150570	Attribut numinuaria DIVA sequence	4.0
1500	D34630	Arabidopsis thaliana mRNA for acetyl-CoA carboxylase, complete cds	0.22
1000	201000	Homo sapiens chromosome 5, P1 clone 1209C1 (LBNL H104),	
1501	AC004498	complete sequence	3E-13
1001	1100011110	Dictyostelium discoideum prespore-specific protein (pspC) gene,	- DE 10
1502	AF104350	partial cds; and unknown gene	0.017
1503	XM 028059	Homo sapiens KIAA1681 protein (KIAA1681), mRNA	0.35
		Mus musculus serotonin N-acetyltransferase (AANAT) gene,	
1504	U83462	complete cds	0.017
1505	AD001534	Borrelia afzelli (isolate UMO1), sequence fragment	0.7
		Rivulus haraldsiolii tRNA-Ile, tRNA-Gln, and tRNA-Met genes,	
		complete sequence; NADH dehydrogenase subunit II gene, complete	
1		cds; tRNA-Trp, tRNA-Ala, tRNA-Asn, tRNA-Cys, and tRNA-Tyr	
		genes, complete sequence; and cytochrome c oxidase subunit I gene.	
1506	AF092379	pa>	0.002
		Homo sapiens clone 11 pur alpha-associated ribosomal RNA gene,	
1507	AF397158	partial sequence	1E-39
		Arabidopsis thaliana putative ribosomal protein (AT4g16720) mRNA,	
1509	AF370352	complete cds	1.2

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
		HIV-1 isolate 97CDKFE4 from Republic of the Congo gag protein	
1510	AF286240	(gag) and pol protein (pol) genes, partial cds	1.2
1511	L29256	Human dinucleotide repeat polymorphism at the DXS1690 locus	5E-50
1512	AF214115	Peromyscus maniculatus H19 mRNA, complete cds	2.1
1513	XM_003799	Homo sapiens protocadherin beta 10 (PCDHB10), mRNA	0.57
1514	AF387030	Homo sapiens DNA polymerase epsilon subunit B (POLE2) gene, exons 14 and 15	0.72
1515	M27054	Homo sapiens asparagine synthetase (TS11) gene, exons 1-3 and promoter region	0.2
		Homo sapiens mesoderm development candidate 2 (MESDC2),	
1516	XM_051852	mRNA	5.9
		Homo sapiens heterogeneous nuclear protein similar to rat helix	
1517	XM_052004	destabilizing protein (FBRNP), mRNA	0.00009
		Homo sapiens hypothetical gene supported by L27560 (LOC92701).	
1518	XM 046730	mRNA	0.53
1519	AF136179	Mus musculus spermine synthase gene, partial cds	0.063
		Onchocerca volvulus nuclear hormone receptor-like peptide, complete	
1520	U19360	cds	1.6
1521	AY039907	Arabidopsis thaliana AT5g41810/K16L22_9 mRNA, complete cds	5.9
		Plasmodium falciparum DNA for flavoprotein subunit of succinate	
1522	D86573	dehydrogenase, complete cds	0.009
1523	NC_002548	Acute bee paralysis virus, complete genome	0.07
1524	U33959	Macaca fascicularis fertilin beta mRNA, complete cds	0,3
		Staphylococcus epidermidis strain SR1 clone step.1043h07 genomic	
1525	AF270048	sequence	. 0.76
		Staphylococcus aureus DNA, complete structure of cassette	
1526	AB047239	chromosome(SCC)-like element, strain:ATCC25923	0.72
1527	U39684	Mycoplasma genitalium section 6 of 51 of the complete genome	0.21
1528	XM_030430	Homo sapiens hypothetical protein (FLJ11127), mRNA	2
1529	D38148	Rotavirus strain A5-13 NSP1 gene, complete cds	0.68
		Mus musculus adult retina cDNA, RIKEN full-length enriched	
1530	AK020846	library, clone:A930011E06, full insert sequence	0.2
1531	AF130049	Homo sapiens clone FLB3411 PRO0852 mRNA, complete cds	0.067
1532	AF332140	Mus musculus c-Fos (Fos) gene, 5' flanking and promoter regions	1.8
		Jasminum odoratissimum tRNA-Leu (trnL) gene, partial intron and 3'	
		exon, and trnL-trnF spacer region; chloroplast gene for chloroplast	
1533	AF231842	product	1.3
1534	NC_001560	Vesicular stomatitis virus, complete genome	0.35
1535	AF213897	Homo sapiens cAMP responsive element modulator (CREM) gene, exon C	0.24
1536	XM_049251	Homo sapiens KIAA0903 protein (KIAA0903), mRNA	0.22
1505		Homo sapiens cDNA FLJ14807 fis, clone NT2RP4001760, weakly similar to PUTATIVE RHO/RAC GUANINE NUCLEOTIDE	
1537	AK027713	EXCHANGE FACTOR	1E-36

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SEQ ID		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
NO	ACCESSN	DESCRIP	P VALUE
		Chloris gayana phosphoenolpyruvate carboxylase (ppc) mRNA,	
1538	AF268091	complete cds	6.1
1539	AC023351	Homo sapiens clone RP11-637J20, complete sequence	1E-49
1540	NC_002161	Bovine parainfluenza virus 3, complete genome	6.4
1541	AE001092	Archaeoglobus fulgidus section 15 of 172 of the complete genome	0.15
		Drosophila melanogaster glutamate receptor DGluRIIB mRNA,	
1542	AF044202	complete cds	0.008
1543	Z72767	S.cerevisiae chromosome VII reading frame ORF YGL245w	1.8
1544	AJ277899	Nicotiana tabacum drepp4 gene, exons 1-4	0.24
1545	XM_005131	Homo sapiens chromatin accessibility complex 1 (CHRAC1), mRNA	0.71
1546	AF270048	Staphylococcus epidermidis strain SR1 clone step.1043h07 genomic sequence	0.67
1547	AL112743	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen deprivation	0.076
		YAP2=AP-1-like stress-induced transcriptional activator	
1548	S68847	[Saccharomyces cerevisiae, W303, Genomic, 2181 nt]	0.21
1549 1551	AF176634 X79988	Reldia sp. Amaya and Smith 578 NADH dehydrogenase subunit F (ndhF) gene, partial cds; chloroplast gene for chloroplast product H.sapiens HIP gene, exon 1	0.54
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	0.21
1552	AK015054	library, clone:4930402M22, full insert sequence	2.2
1553	U07731	Human quinone oxidoreductase2 (NQO2) gene, introns 1 and 2 and exon 2	0.42
1554	AE001370	Plasmodium falciparum chromosome 2, section 7 of 73 of the complete sequence	0.02
1555	AF222996	Danio rerio nocA-like Zn-finger protein (nlz) mRNA, complete cds	0.59
1556	AJ000080	Trypanosoma brucei hsp100 and gpi-plc genes	0.00004
1557	D30745	Xenopus Iaevis MRP RNA gene	5
1558	X17256	Yeast THR4 gene for threonine synthase (EC 4.2.99.2)	0.64

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Table 3A Nearest Neighbor (BlastN vs. Genbank) SEQ ID NO ACCESSN DESCRIP P VALUE 1568 Z29373 H.sapiens gene for neural cell adhesion molecule L1 1569 AF138876 Bacillus thuringiensis isolate M15 insertion sequence IS231N 0.48 1570 AF043087 Hordeum vulgare dehydrin 1 (dhn1) gene, complete cds 2.1 AE002293 1571 0.57 Chlamydia muridarum, section 25 of 85 of the complete genome Rattus norvegicus gene for hepatocyte nuclear factor 3 gamma, partial 1572 AB017043 3.7 1573 XM_038951 Homo sapiens hypothetical protein MGC15875 (MGC15875), mRNA 0.43 Mus musculus galectin-11 mRNA, variant b, complete cds. 1574 AF244979 alternatively spliced 0.24 1575 XM 006069 Homo sapiens wee1+ (S. pombe) homolog (WEE1), mRNA 6E-10 1576 AK025037 Homo sapiens cDNA: FLJ21384 fis, clone COL03354 1.7 S. cerevisiae chromosome XIV reading frame ORF YNL167c 1577 Z71443 Arabidopsis thaliana protein synthesis initiation factor 4G (EIF4G) 1578 AF263518 gene, complete cds 0.59 AE000680 1579 Aquifex acolicus section 12 of 109 of the complete genome 1.2 Plasmodium falciparum chromosome 2, section 41 of 73 of the 1580 AE001404 complete sequence 1.5 Candida albicans partial mRNA for serine/threonine kinase (sha3) 1581 AJ390510 0.18 1582 AC004173 Homo sapiens clone UWGC:y23x011 from 6p21, complete sequence 0.002 1583 Z71687 S.cerevisiae chromosome XIV reading frame ORF YNR072w 2.1 1585 X95065 H.sapiens dinucleotide repeat (GT) from chromosome 13q (229bp) 0.18 H. sapiens (D10S593) DNA segment containing (CA) repeat; clone 1586 Z24310 AFM311vb1; single read 0.002 Trichophyton rubrum mitochondrial cytb gene and NADH1 to 1587 Y18476 NADH5 genes 6.9 1588 Z47046 Human cosmid QLL2C9 from Xq28 0.00007 1589 Z71687 S.cerevisiae chromosome XIV reading frame ORF YNR072w 1.6 1590 XM 005485 Homo sapiens KIAA0414 protein (KIAA0414), mRNA 0.002 NM 024364 1591 Rattus norvegicus hairless (hr), mRNA 0.22 Caenorhabditis elegans homogentisate 1,2-dioxygenase (hgo) gene, 1593 AF136150 complete cds 1.6 1594 XM 033896 Homo sapiens insulin-like growth factor 2 receptor (IGF2R), mRNA 1.4 1595 AJ404228 Saccharomyces douglasii mitochondrial tRNA gene cluster 0.008 Homo sapiens, acid phosphatase, prostate, clone MGC:12236 1596 BC007460 0.029 IMAGE:3951204, mRNA, complete cds Mus musculus RIKEN cDNA 0610011N22 gene (0610011N22Rik). 1597 NM 024201 5.3 1598 U88166 Caenorhabditis elegans cosmid M01A12 0.61 Plasmodium falciparum chromosome 2, section 31 of 73 of the 1599 AE001394 complete sequence 2

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Corynebacterium glutamicum partial nrdF gene for ribonucleotide	
		reductase beta-subunit and ctaD gene for cytochrome aa3 oxidase	1
1600	AJ306417	subunit I	1.9
1601	AF287270	Homo sapiens mucolipin (MCOLN1) gene, complete cds	6
		Chlamydophila pneumoniae AR39, section 37 of 94 of the complete	
1602	AE002207	genome	0.083
		Mus musculus 18 days embryo cDNA, RIKEN full-length enriched	
1603	AK003449	library, clone:1110004P21, full insert sequence	0.2
		Arabidopsis thaliana AtERF-1 mRNA for ethylene responsive element	
1604	AB008103	binding factor 1, complete cds	0.79
		Lactococcus lactis subsp. lactis IL1403 section 168 of 218 of the	
1605	AE006406	complete genome	0.18
		Homo sapiens, clone MGC:18053 IMAGE:4148889, mRNA,	
1606	BC008122	complete cds	1.5
		Chicken mRNA for c-maf proto-oncogene product c-Maf long form,	
1607	D28598	complete cds	0.44
1608	X97314	M.sativa mRNA for cdc2 kinase homologue, cdc2MsC	1.5
	-	Homo sapiens genomic DNA, chromosome 21q22.2, clone:10C6,	
1609	AP001407	LB7T-ERG region, complete sequence	1.1
		Homo sapiens similar to heterogeneous nuclear ribonucleoprotein L	
1611	XM_017891	(H. sapiens) (LOC91538), mRNA	1.4
1		Corrigiola littoralis ORF2280 gene homolog, chloroplast gene	
1612	U48562	encoding chloroplast protein, partial cds	0.074
		Homo sapiens similar to tumor protein p53-binding protein (H.	
1613	XM_036274	sapiens) (LOC65591), mRNA	2
i l			
1614	AJ271039	Plasmodium yoelii yoelii partial mrk gene for MO15-related kinase	0.001
1615	AB016467	Datisca cannabina chloroplast gene for maturase K, partial cds	0.25
1616	AF027807	Homo sapiens beta-casein (CSN2) gene, complete cds	0.074
		Mus musculus adult male kidney cDNA, RIKEN full-length enriched	
1617	AK002822	library, clone:0610039A20, full insert sequence	0.55
			l
1618	U67580	Methanococcus jannaschii section 122 of 150 of the complete genome	0.032
]			
1619	U89140	Mus musculus aldose reductase gene, promoter region and exon 1	0.48
		Drosophila melanogaster genomic scaffold 142000013385566,	
1620	AE002799	complete sequence	0.17
1621	L14855	Bos taurus neurexin I-alpha mRNA, complete cds	0.48
1622	AF379854	Cloning vector pVLH/hsp, complete sequence	2
		S.tuberosum U1snRNA variant genes U1-1, U1-2, U1-3, U1-4, U1-5	1
1623	Z11883	and U1-6	0.54
		Mus musculus ILTIFa gene for IL-TIF alpha protein (IL-22), exons 1a	
1624	AJ294727	5	0.011
		Homo sapiens hypothetical protein DKFZp434A1319	
1625	XM_043073	(DKFZP434A1319), mRNA	1.9

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	!	Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		*	
NO	ACCESSN	DESCRIP	P VALUE
1626	M84732	Plasmodium yoelii sporozoite surface protein 2 gene, complete cds	0.007
		G.hirsutum late embryogenesis-abundant protein 2-D (LeaA2-D)	
1627	M83304	mRNA, complete cds	9.8
1628	U32725	Haemophilus influenzae Rd section 40 of 163 of the complete genome	0.14
		Homo sapiens cytochrome P450 3A4 (CYP3A4) gene, promoter	
1629	AF185589	region	0.027
		Human DNA sequence from clone RP11-389N9 on chromosome 6,	
1630	AL591667	complete sequence [Homo sapiens]	0.15
1631	AB058397	Oryza sativa CHS gene for chalcone synthase, complete cds	0.16
_1632	M11043	Mouse DNA fragment that hybridizes to HSV-1 Smal A fragment	1.5
		Staphylococcus aureus putative membrane protein MprF (mprF) gene,	
1633	AF145699	complete cds; and unknown gene	1.5
1634	AF088902	Mus musculus SLIT1 protein (Slit1) mRNA, partial cds	0.5
1635	AB048882	Macaca fascicularis brain cDNA, clone:QnpA-16525	0.21
1636	X98369	M.musculus mRNA for SOX15 protein	0.24
1637	AF131767	Homo sapiens clone 24900 mRNA sequence	0.25
		Homo sapiens mRNA; cDNA DKFZp586F1924 (from clone	
1638	AL117657	DKFZp586F1924)	1.9
1639	AB042973	Nicotiana tabacum wizz gene, 5' flanking region, partial cds	0.072
1640	AJ272083	Staphylococcus aureus empbp gene for extracellular matrix and plasma binding protein, strain SA 113	0.000
1040	AJZ12063	Homo sapiens, clone MGC:15151 IMAGE:3538202, mRNA,	0.008
1641	BC009423	complete cds	1.9
1041	BC009423	Saccharomyces sp. CID1 ATP synthase subunit 8 (ATP8) gene,	1.9
1642	AF114905	mitochondrial gene encoding mitochondrial protein, complete cds	0.18
1042	711 114705	Staphylococcus epidermidis strain SR1 clone step. 4052d03a genomic	0.10
1643	AF270417	sequence	0.063
1644	X91340	G.hybrida chs-like gene	0.51
1044	10 1540	Homo sapiens autosomal highly conserved protein (AHCP) gene,	0.51
1645	AF097026	complete cds	0.008
1646	XM 028810	Homo sapiens 67292 (KIAA1755), mRNA	5,5
		Homo sapiens mRNA; cDNA DKFZp564P046 (from clone	
1647	AL049339	DKFZp564P046)	2.1
		Plasmodium falciparum chromosome 2, section 35 of 73 of the	
1648	AE001398	complete sequence	0.13
1649	M55171	Mouse opsin (MOPS) gene, complete cds	0.52
		Arabidopsis thaliana AtERF-1 mRNA for ethylene responsive element	
1650	AB008103	binding factor 1, complete cds	0.66
		Human DNA sequence from clone RP11-64M7 on chromosome 6,	
1651	AL589705	complete sequence [Homo sapiens]	0.5
1652	X79559	T.trypanoides (R1) 18S rRNA gene	0.069
1653	XM_048416	Homo sapiens PRO0478 protein (PRO0478), mRNA	0.0001

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Dasyurus maculatus isolate QM.E1 mitochondrial D-loop, partial	
1654	AF082761	sequence	0.063
1655	AF282568	Homo sapiens clone 20ptel_c15bt7 sequence	4E-39
		Pyrus pyrifolia gene for UDP-glucose pyrophosphorylase, complete	
1656	AB027617	cds	0.19
1657	AK023864	Homo sapiens cDNA FLJ13802 fis, clone THYRO1000186	0.2
1658	NM_009905	Mus musculus CDC-like kinase (Clk), mRNA	1.7
1659	AL023846	Caenorhabditis elegans cosmid Y52B11C, complete sequence	2.3
1660	AF334780	Homo sapiens pregnancy-induced growth inhibitor OKL38 gene, partial cds	0.73
		Homo sapiens mitogen-activated protein kinase kinase kinase 1	
1661	XM_042066	(MAP3K1), mRNA	1.9
1662	XM_034872	Homo sapiens KIAA0202 protein (KIAA0202), mRNA	0.23
		Homo sapiens (subclone 1_g7 from BAC H76) DNA sequence,	
1663	AC002252	complete sequence	3E-14
1664	AK023113	Homo sapiens cDNA FLJ13051 fis, clone NT2RP3001447	2E-20
1665	BC003552	Homo sapiens, calnexin, clone MGC:1196 IMAGE:3546389, mRNA, complete cds	0.076
1000	AT007142	Tapinaspis sp. 12S small subunit ribosomal RNA gene, partial	
1666	AF097142	sequence; mitochondrial gene for mitochondrial product	1.5
1667	47 117100	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen	١
1668	AL117103 NM_013825	deprivation Mus musculus lymphocyte antigen 75 (Ly75), mRNA	6.1
1669	NM_013823 NM_007043	Homo sapiens HIV-1 rev binding protein 2 (HRB2), mRNA	0.74
1670	Z27359		7E-22
1671	AK021535	G.hirsutum (AD52) copia-like reverse transcriptase fragments	0.24
10/1	AK021333	Homo sapiens cDNA FLJ11473 fis, clone HEMBA1001712	0.25
1672	Y00145	Dictyostelium discoideum UDP glucose pyrophosphorylase gene (UDPGP; EC 2.7.7.9)	
1673	M63291	Drosophila silvestris alcohol dehydrogenase gene, complete cds	0.22
10/3	10103291	Diosophila silvestris alcohol denydrogenase gene, complete cus	0.028
		I-G. D. D. D. C. 1. 17/01	
1674	AF100401	Influenza B virus B/Guangdong/5/94 segment 8 nonstructural protein (NS1) and nonstructural protein (NS2) genes, complete cds	
1675	U80449	Caenorhabditis elegans cosmid R10A10	0.2
1676	X15634	C. elegans core histone genes H4(his-10) and H3(his-9)	0.022
1677	XM 050366	Homo sapiens zinc finger protein, X-linked (ZFX), mRNA	0.022
10//	AM_030300		0.23
1678	AE007046	Mycobacterium tuberculosis CDC1551, section 132 of 280 of the	0.000
10/6	AE007040	complete genome Homo sapiens glucose-6-phosphatase, catalytic (glycogen storage	0.062
1679	XM 008207	disease type I, von Gierke disease) (G6PC), mRNA	477.10
10/2	ZLVI_000207	Hemicentrotus pulcherrimus mRNA for transcription factor HpLim1.	4E-18
1680	AB049118	complete cds	0.21
1000	ADV+9110	lounding cris	0.21
1681	XM 002421	Homo sapiens hypothetical protein FLJ20254 (FLJ20254), mRNA	4.1
1682	X96783	H. sapiens Syt V gene (genomic and cDNA sequence)	0.000004
	5765		

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
		Plasmodium berghei extrachromosomal plastid PB-2, tRNA-Pro,	1 TILLOW
		tRNA-Gin, tRNA-Lys, tRNA-Asp, tRNA-Ser, tRNA-Tyr, tRNA-Met.	
1		tRNA-Leu, tRNA-Cys, and tRNA-His genes, complete sequence, rps4	ľ
		gene, complete cds, tRNA-Thr gene, complete sequence, and large	
1683	U79732	subu>	0.001
1684	NC_002619	Pteropus scapulatus mitochondrion, complete genome	6.3
1685	L16679	Caenorhabditis elegans cosmid K07D8, complete sequence	1.8
1686	Y08924	P.falciparum mRNA for AARP2 protein	0.003
1687	U32760	Haemophilus influenzae Rd section 75 of 163 of the complete genome	2.2
1688	AF298624	Dictyostelium discoideum chromosome 2 repeat region	0.009
1689	XM_031155	Homo sapiens RAS protein activator like 2 (RASAL2), mRNA	0.33
1690	AE006899	Sulfolobus solfataricus section 258 of 272 of the complete genome	5
1691	U67556	Methanococcus jannaschii section 98 of 150 of the complete genome	0.42
		Mus musculus 10, 11 days embryo cDNA, RIKEN full-length	
1692	AK012631	enriched library, clone:2810001G20, full insert sequence	0.064
		Homo sapiens, clone MGC:16308 IMAGE:3836116, mRNA,	
1693	BC007399	complete cds	0.022
		Human DNA sequence from clone RP11-110C10 on chromosome 20.	
1694	AL389897	Contains GSSs, complete sequence [Homo sapiens]	1E-26
		H.sapiens CpG island DNA genomic Mse1 fragment, clone 68h2,	
1695	Z62421	forward read cpg68h2.ft1a	4E-70
1696	BC010535	Homo sapiens, clone IMAGE:3456494, mRNA	0.002
1697	Z68319	Caenorhabditis elegans cosmid T23G7, complete sequence	0.18
		Mus musculus transcriptional activator alpha-NAC (Naca) gene,	
1698	U48363	complete cds .	0.058
		Ustilago nuda 18S ribosomal RNA gene, partial sequence; internal	
		transcribed spacer 1, 5.8S ribosomal RNA gene and internal	
		transcribed spacer 2, complete sequence; and 28S ribosomal RNA	
1699	AF135430	gene, partial sequence	1.4
		Homo sapiens similar to enhancer of rudimentary (Drosophila)	
1701	XM_007461	homolog (H. sapiens) (LOC65719), mRNA	0.18
1702	NM_025083	Homo sapiens hypothetical protein FLJ21128 (FLJ21128), mRNA	e-124
		Homo sapiens uncharacterized hematopoietic stem/progenitor cells	
1703	XM_050592	protein MDS032 (MDS032), mRNA	0,068
		Homo sapiens Makorin RING zinc-finger protein 2 (MKRN2) gene,	
1704	AF277167	exon 4	2
		Plasmodium falciparum chromosome 2, section 70 of 73 of the	
1705	AE001433	complete sequence	0.67
1706	AF075003	Homo sapiens full length insert cDNA YH97B03	0.001
		Mus musculus transcriptional regulator, SIN3A (yeast) (Sin3a),	
1707	NM_011378	mRNA	0.62

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NO ACCESSN DESCRIP P VALUE			Table 3A Nearest Neighbor (BlastN vs. Genbank)	
AE001321 Chlamydia trachomatis section 48 of 87 of the complete genome 0.026				
Homo sapiens similar to elongation of very long chain fatty acids (FEN.IEEOZ, SUR4/Elo3), yeast)-like 1 (H. sapiens) (L.OC92652), mm/s (FEN.IEEOZ, SUR4/Elo3), years)-like 1 (H. sapiens) (L.OC92652), mm/s (H. sapiens) (H. sapiens) (L.OC92652), mm/s (H. sapiens) (H				
CENIL/EioZ, SUR4/Eio3, yeasty-like 1 (H. sapiens) (LOC92652), mRNA	1708	AE001321		0.026
1709 XM_046388 mRNA 6.1				
MOR6.5=ouabain resistance gene {repeat sequence} { mice, Genomic, 693 nt] MOR6.5=ouabain resistance gene {repeat sequence} { mice, Genomic, 693 nt] 1711				ļ
1710	1709	XM_046388		6.1
1711 AB055378 Macaca fascicularis brain cDNA, clone:QfIA-13294 0.22	1510	055505		
1712 XM_043597 Homo sapiens centaurin-alpha 2 protein (HSA272195), mRNA 1E-20				
AE007426 Streptococcus pneumoniae section 109 of 194 of the complete genome 0.32				
Miss musculus adult male testis cDNA, RIKEN full-length enriched library, clone;4932411G06, full insert sequence e-129	1/12	AIVI_043397	rionio sapiens centaurin-aipna 2 protein (HSA272195), mRNA	1E-20
Miss musculus adult male testis cDNA, RIKEN full-length enriched library, clone;4932411G06, full insert sequence e-129	1712	AE007426	Strentosocous manumonica socion 100 of 104 of the complete consum	0.22
1714	1/13	AE007420		0.32
1715 AL122033 S. pombe chromosome II cosmid c1921 0.53	1714	AK016519		0.120
Mus musculus 10, 11 days embryo cDNA, RIKEN full-length enriched library, clone:2810401603, full insert sequence 0.075				
AK012960	1110	12010000		0.55
Human DNA sequence from clone RP1-473/14 on chromosome X, complete sequence [Homo sapiens] 0.19	1716	AK012960		0.075
1718 AL590123 complete sequence [Homo sapiens] 0.19 1718 D17343 Ratius norvegicus cytochrome P450 2B15 gene, exon 1 0.66 1719 XM_028703 Homo sapiens KIAA0889 protein (KIAA0889), mRNA 0.00004 1720 AB060865 sequence 1720 AB060865 sequence 2.3 1721 AL133397 Latinity 1720 Latinity				0.072
1718 D17343 Rattus norvegicus cytochrome P450 2B15 gene, exon 1 0.66	1717	AL590123		0.19
1719 XM_028703 Homo sapiens KIAA0889 protein (KIAA0889), mRNA 0.00004	1718	D17343		
1720	1719	XM_028703		0.00004
Human DNA sequence from clone RP1-213IIp on chromosome 22q11.21-12.2, complete sequence [Homo sapiens] 0,083			Macaca fascicularis brain cDNA clone:QtrA-11953, full insert	
1721	1720	AB060865	sequence	2.3
Mitochondrion Drosophila eugracilis ND2 and COI genes (partial) and genes for RNA-Trp, RNA-Tyr, and RNA-Cys 0.074			Human DNA sequence from clone RP1-213J1p on chromosome	
1722 X38913 and genes for tRNA-Trp, tRNA-Trp, tRNA-Trp, tand tRNA-Cys 0.074	1721	AL133397		0,083
Homo sapiens Xp22 Cosmid U239B3 (from Lawrence Livermore X bibrary) complete sequence 0.0008				
1723	1722	X58913		0.074
Lactococcus lactis subsp. lactis IL1403 section 84 of 218 of the complete genome 1.8				
1724 AE006322 complete genome 1.8 1725 Z71290 S.cerevisiae chromosome XIV reading frame ORF YNL014w 0.67 1726 XM_005723 Homo sapiens hypothetical protein PRO2859 (PRO2859), mRNA 0.014 1727 AF123488 South River virus polyprotein mRNA, complete cds 0.57 1728 D78176 Mouse DNA for primary transcript of atrial natriuretic clearance receptor 0.061 1730 XM_006871 Homo sapiens sethanolamine kinase (EKID), mRNA 4.1 1731 AL078640 Contains an EST, complete sequence [Homo sapiens] 0.07 1732 AC092030 Homo sapiens clone RFI1-636112, complete sequence 0.029 Mus musculus 18 days embryo cDNA, RIKEN full-length enriched library, clone:1110003P05, full insert sequence 0.16 1734 AF131767 Homo sapiens clone Ry900 mRNA sequence 0.2 1735 Z74246 Scerevisiae chromosome IV reading frame ORF YDL198c 0.023 1736 NM_021401 Mus musculus secreted and transmembrane I (sectm1), mRNA 0.069	1723	AC002359		0.0008
1725 Z71290 S.cerevisiae chromosome XIV reading frame ORF YNL014w 0.67				
1726 XM_005723 Homo sapiens hypothetical protein PRO2859 (PRO2859), mRNA 0.014 1727				
1727 AF123488 South River virus polyprotein mRNA, complete cds 0.57	1725	Z71290	S.cerevisiae chromosome XIV reading frame ORF YNL014w	0.67
1727 AF123488 South River virus polyprotein mRNA, complete cds 0.57		77 - 00-500		
1728 D78176 Mouse DNA for primary transcript of atrial natriuretic clearance receptor receptor 0.061				
1728 D78176 receptor 0.061 1730 XM_006871 Homo sapiens ethanolamine kinase (EKII), mRNA 4.1 1731 AL078640 Human DNA sequence from clone 536P6 on chromosome 22. 0.07 1732 AC092030 Homo sapiens clone RP11-636H2, complete sequence 0.029 Mus musculus 18 days embryo cDNA, RKEM full-length enriched library, clone:1110003P05, full insert sequence 0.16 1734 AF131767 Homo sapiens clone 24990 mRNA sequence 0.2 1735 Z74246 S.cerevisiae chromosome IV reading frame ORF YDL198c 0.023 1736 NM_021401 Mus musculus secreted and transmembrane 1 (Sectm1), mRNA 0.069 Buchnera aphidicola DNA polymerase III beta subunit (dmkN) gene, Buchnera aphidicola DNA polymerase III beta subunit (dmkN) gene,	1/2/	AF123488		0.57
1730 XM_006871 Homo sapiens ethanolamine kinase (EKII), mRNA 4.1 1731 AL078640 Contains an EST, complete sequence (Flomo sapiens) 0.07 1732 AC092030 Homo sapiens clone RP11-636112, complete sequence 0.029 Mus musculus 18 days embryo cDNA, RIKEN full-length enriched birary, clone: 1110003F05, full insert sequence 0.16 1734 AF131767 Homo sapiens clone 24900 mRNA sequence 0.2 1735 Z74246 S.cerevisiae chromosome IV reading frame ORF YDL198c 0.023 1736 NM_021401 Mis musculus secreted and transmembrane I (Sectm1), mRNA 0.069 Buchnera aphidicola DNA polymerase III beta subunit (dnaN) gene, Buchnera sphidicola DNA polymerase III beta subunit (dnaN) gene,	1700	D79176		0.061
Human DNA sequence from clone 536P6 on chromosome 22.				
1731 AL078640 Contains an EST, complete sequence [Homo sapiens] 0.07 1732 AC092030 Homo sapiens clone RPI1-636112, complete sequence 0.029 Mus musculus 18 days embryo cDNA, RIKEBY full-length enriched 0.16 1733 AK003349 library, clone: 1110003P05, full insert sequence 0.16 1734 AF131767 Homo sapiens clone 2990 mRNA sequenco 0.2 1735 Z74246 S. cerevisiae chromosome IV reading frame ORF YDL198c 0.023 1736 NM 021401 Mus musculus secreted and transmembrane 1 (Sectm1), mRNA 0.069 Bucharea aphidicola DNA polymenase III beta subunit (dnaN) gene, Bucharea aphidicola DNA polymenase III beta subunit (dnaN) gene,	1750	2111_000071		7.1
1732 AC092030 Homo sapiens clone RP11-636112, complete sequence 0.029 Mus musculus 18 days embryo cDNA, RIKEN full-length enriched 0.16 1733 AK003349 bibrary, clone: 111003078, full insert sequence 0.16 1734 AF131767 Homo sapiens clone 24900 mRNA sequence 0.2 1735 Z74246 S.cerevisiae chromosome IV reading frame ORF YDL198c 0.023 1736 NM_021401 Mus musculus secreted and transmembrane I (Sectm1), mRNA 0.069 Buchnera aphidicola DNA polymerase III beta subunit (dnaN) gene, Buchnera aphidicola DNA polymerase III beta subunit (dnaN) gene,	1731	AT 078640		0.07
Mins musculus 18 days embryo cDNA, RIKEN full-length enriched library, clone: 1110003F05, dail insert sequence 0.16 1734 AF131767 Home sapiens clone 24900 mRNA sequence 0.2 1735 Z74246 S. cerevisiae chromosome IV reading frame ORF YDL198c 0.023 1736 NM_021401 Mus musculus secreted and transmembrane I (Sectm1), mRNA Bucharea aphidicola DNA polymerase III beta subunit (dnaN) gene,				
1733 AK003349 library, clone:1110003F05, full insert sequence 0.16 1734 AF131767 Homo sapiens clone 24900 mRNA sequence 0.2 1735 Z74246 S. cerevisiae chromosome IV reading frame ORF YDL198c 0.023 1736 NM_021401 Mus musculus secreted and transmembrane 1 (Sectm1), mRNA 0.069 Buchnera aphidicola DNA polymerase III beta subunit (dnaN) gene, 10.000				
1734 AF131767 Homo sapiens clone 24900 mRNA sequence 0.2 1735 Z74246 S.cerevisiae chromosome IV reading frame ORF YDL198c 0.023 1736 NM_021401 Mus musculus secreted and transmembrane I (Sectm1), mRNA 0.069 Buchnera aphidicola DNA polymerase III beta subunit (dnaN) gene, 0.069	1733	AK003349		0.16
1736 NM 021401 Mus musculus secreted and transmembrane I (Sectru1), mRNA 0.069 Buchnera aphidicola DNA polymerase III beta subunit (dnaN) gene,	1734	AF131767		
1736 NM_021401 Mus musculus secreted and transmembrane 1 (Sectm1), mRNA 0.069 Buchnera aphidicola DNA polymerase III beta subunit (dnaN) gene,	1735	Z74246		0.023
Buchnera aphidicola DNA polymerase III beta subunit (dnaN) gene,	1736	NM_021401		0.069
1737 AF213931 complete cds 0.68			Buchnera aphidicola DNA polymerase III beta subunit (dnaN) gene,	
	1737	AF213931	complete cds	0.68

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WC0214500 [fle://E/WO0214500 opc]__

		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Homo sapiens clone 20r DH4-17/BCL-1 gene fusion reciprocal	
1738	AF288892	breakpoint sequence	2
1739	X99719	S.enterica hsdM, hsdS & hsdR genes	0.23
1740	XM_041866	Homo sapiens KIAA1468 protein (KIAA1468), mRNA	5.9
1741	AY010113	Homo sapiens unknown mRNA sequence	0.64
l		Mus musculus adult male tongue cDNA, RIKEN full-length enriched	
1742	AK009187	library, clone:2310006I24, full insert sequence	4E-23
1743	AF130074	Homo sapiens clone FLB9348 PRO2523 mRNA, complete cds	0.14
1744	AJ249165	Plasmodium falciparum gc-beta gene, exons 1-13	0.002
		Mus musculus 0 day neonate skin cDNA, RIKEN full-length enriched	Ì
1745	AK014565	library, clone:4632411B12, full insert sequence	0,54
1746	M15058	Vaccinia virus (strain WR) HindIII D fragment DNA, complete	0,15
		M.pulmonis DNA for insertion sequence IS1138 encoding putative	
1747	Z16416	transposase	0.026
1748	XM_047638	Homo sapiens Fzr1 protein (FZR1), mRNA	1.2
	*****	Anabaena sp. phycobilisome core component (apcF) gene, complete	
1750	U21853	cds, and glutamine synthetase (glnA) gene, partial cds	0.076
1751	AF169454	Neisseria meningitidis strain Z2491 clone Cm045 unknown sequence	0.19
		Apanteles canarsiae 16S large subunit ribosomal RNA gene, partial	
1752	AF102750	sequence; mitochondrial gene for mitochondrial product	0,066
		Plasmodium falciparum sexual stage antigen (s16) gene, promoter and	
1753	AF034389	partial cds	0.18
		Staphylococcus aureus DNA, complete structure of cassette	
1754	AB047239	chromosome(SCC)-like element, strain:ATCC25923	1.5
1755	X98669	A.thaliana zat1 gene	0,023
1756	X16619	Chlamydomonas reinhardtii ARG7 gene for argininosuccinate lyase	1.5
1757	XM_005488	Homo sapiens KIAA0354 gene product (KIAA0354), mRNA	0.063
1758	AF110966	HIV-1 isolate C-96BW04.10 country Botswana, complete genome	1.1
1759	AF247193	Mus musculus endobrevin (Vamp8) gene, exon 1	5.4
l I		Homo sapiens proteasome (prosome, macropain) 26S subunit, non-	
1760	XM_045302	ATPase, 5 (PSMD5), mRNA	0.0008
1		Lactococcus lactis subsp. lactis IL1403 section 112 of 218 of the	
1761	AE006350	complete genome	0.69
		Anguilla anguilla mitochondrial DNA for partial tRNA-Thr and	
1762	AJ247001	tRNA-Pro genes and D-loop, isolate RM11	0.052
		Cicer arietinum mRNA for rac-type small GTP-binding protein,	
1763	AB024996	complete cds	1.6
1764	M13945	Mus musculus pim-1 protein kinase gene, complete eds	5.7
,,,,	17740001		
1765	AF106842	Platanus racemosa PlaraLFY protein (PlaraLFY) gene, complete cds	0.22
1200	ATO(0000	Staphylococcus epidermidis strain SR1 clone step.1024f03 genomic	
1766	AF269822	sequence	7.5
1767	AF256830	Salmo salar clone BHMS549 microsatellite sequence	1.8

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
1768	AE007395	Streptococcus pneumoniae section 78 of 194 of the complete genome	1.6
1769	AB028896	Streptococcus agalactiae type-specific capsular polysaccharide biosynthesis gene region (cpsIaA to ung gene), complete cds	0.025
1770	AL583831	Human DNA sequence from clone RP11-469L10 on chromosome 6, complete sequence [Homo sapiens]	0,15
1771	AL354683	Human DNA sequence from clone RP11-259K11 on chromosome 20, complete sequence [Homo sapiens]	0.21
1772	XM_047524	Homo sapiens procollagen (type III) N-endopeptidase (PCOLN3), mRNA	4.6
1773	AC084500	Caenorhabditis briggsae cosmid G13G15, complete sequence	0.076
1774	M13945	Mus musculus pim-1 protein kinase gene, complete cds	6.1
1775	L27474	Human arginase (ARGI) gene dinucleotide repeat polymorphism, intron 5	0.21
1776	Z79696	Caenorhabditis elegans cosmid F54F3, complete sequence	1.9
1777	AY018282	Oryza sativa microsatellite MRG0607 containing (AT)X14, closest to marker C132, genomic sequence	0.071
1778	AF114938	Saccharomyces sp. IFO 1815 small subunit ribosomal RNA gene, mitochondrial gene for mitochondrial RNA, complete sequence	0,078
1779	AF170085	Homo sapiens Charcot-Leyden crystal protein gene, partial sequence	6E-18
1780	Y08919	P.flesus mRNA for p53 protein	0.082
1781	L46868	Trypanosoma brucei dominant expression site (DES) gene, promoter region	2.2
1782	AF157008	Prunus dulcis self-incompatibility associated ribonuclease gene, complete cds	0.44
1783	AB017459	Carabus sylvestris mitochondrial ND5 gene, isolate: Baumgartner Austria, partial cds	0.029
1784	AJ294714	Pichia anomala leu2 gene for beta-isopropylmalate dehydrogenase	0.26
1785	AF125972	Caenorhabditis elegans cosmid H25P19	0.64
1786	AB015413	Helicobacter pylori cagA gene, complete cds, strain ATCC43526	1.5
1787	AK021477	Homo sapiens cDNA FLJ11415 fis, clone HEMBA1000942	7E-12
1788	AC006812	Caenorhabditis elegans cosmid Y64H9A, complete sequence	0,63
1789	XM_004181	Homo sapiens megakaryocyte-enhanced gene transcript 1 protein (MEGT1), mRNA	7E-66
1790	AJ236640	Homo sapiens chromosome 22 CpG island DNA, genomic Mse1 fragment, clone 22CGIB49A20 , complete read	9E-19
1791	XM_041751	Homo sapiens discoidin domain receptor family, member 1 (DDR1), mRNA	3.5
1792	AL034546	Human DNA sequence from clone RP5-898I4 on chromosome 22q13.33 Contains a GSS and a putative CpG island, complete sequence [Homo sapiens]	0.000005
1/92	MLU34346	Buchnera aphidicola plasmid pBUs1, repA2 gene, leuA gene and	0,0000005
1793	AJ006873	ORF1	0.23

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
1822	BC003900	Mus musculus, Similar to hypothetical protein 384D8_6, clone MGC:6766 IMAGE:3601298, mRNA, complete cds	0.51
1823	XM_050561	Homo sapiens KIAA0700 protein (KIAA0700), mRNA	0.51
1824	S66407	FLT4=receptor tyrosine kinase isoform FLT4 long {3' region, alternatively spliced} [human, mRNA Partial, 216 nt]	8E-36
1825	AY008805	Equus caballus retinoblastoma (RB1) gene, partial cds	0.058
1826	AJ276629	Rattus norvegicus Sacm21/RT1-A intergenic region, haplotype RT1n and partial RT1-A gene for MHC Class I antigen	3.7
1827	AF029975	Eucalyptus grandis MADS box protein (EGM1) mRNA, complete cds	1.2
1828	AL109708	Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 254679	0.019
1829	AF270353	Staphylococcus epidermidis strain SR1 clone step.4025a02 genomic scquence	0.051
1831	U72236	Dictyostelium discoideum ModA (modA) gene, complete cds	0.5
1832	L15234	Mycoplasma pneumoniae Cys-tRNA, Pro-tRNA, Met-tRNA, Ile- tRNA, Ser-tRNA, fMet-tRNA, Asp-tRNA and Phe-tRNA genes	1.1
1833	XM_012722	Homo sapiens RNA (guanine-7-) methyltransferase (RNMT), mRNA	0.5
1834	AF108228	Ginkgo biloba GinLFY protein (GinLFY) gene, complete cds	4.2
1836	XM_049943	Homo sapiens topoisomerase (DNA) II alpha (170kD) (TOP2A), mRNA	0.47
1837	NC_002674	Aulopus japonicus mitochondrion, complete genome	3.1
1838	AE001399	Plasmodium falciparum chromosome 2, section 36 of 73 of the complete sequence	0,008
1839	U53695	Leishmania enriettii multidrug resistance (lemdr1) gene, 5' UTR	0.5
1840	AF257022	Salmo salar clone BHMS7-009 microsatellite sequence	0.7
1841	AF239571	Acineta superba (RNA-Leu (trnL) gene, partial sequence; and trnL-F intergenic spacer region, complete sequence; chloroplast gene for chloroplast product	5.4
1842	AL032651	Caenorhabditis elegans cosmid Y6D1A, complete sequence	0.019
1844	XM_034229	Homo sapiens similar to hypothetical protein FLJ14260 (H. sapiens) (LOC90803), mRNA	0,068
1845	AL137121	Human DNA sequence from clone RP11-122K21 on chromosome 13, complete sequence [Homo sapiens]	e-147
1846	AF137535	Danio rerio clone Dare-DAXX DAXX protein mRNA, partial cds	1.7
1847	AK023539	Homo sapiens cDNA FLJ13477 fis, clone PLACE1003638	0.21
1848	AF218799	Gallus gallus clone LDH-BT1 lactate dehydrogenase B mRNA, complete cds	6.4
1849	AF250345	Xenopus laevis early growth response protein mRNA, complete cds	0.52
1850	BC006858	Mus musculus, clone IMAGE:3594635, mRNA	1.5
1852	XM_006782	Homo sapiens cyclin T1 (CCNT1), mRNA	0.092
1853	J03883	Rabbit zeta-1 globin processed pseudogene, complete cds	1.5

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			I
NO	ACCESSN	DESCRIP	P VALUE
1854	AF349951	Homo sapiens HP95 mRNA, complete cds	3E-09
		Candidatus Carsonella ruddii natural-host Bactericera cockerelli RNA	
		polymerase beta subunit (rpoB) and RNA polymerase beta-prime	İ
1855	AF268062	subunit (rpoC) genes, partial cds	0.17
		Uncultured gamma proteobacterium SUR-ATT-8 16S ribosomal RNA	
1856	AF114508	gene, complete sequence	1.5
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1857	AE006785	Sulfolobus solfataricus section 144 of 272 of the complete genome	0.091
1858	AF078684	Psoroptes ovis glutathione S-transferase mRNA, complete cds	0.0006
		Mus musculus, FK506 binding protein 1a (12 kDa), clone MGC:6253	
1859	BC004671	IMAGE:3498682, mRNA, complete cds	0.12
1860	NM 025083	Homo sapiens hypothetical protein FLJ21128 (FLJ21128), mRNA	. 101
1800	1NJVI_U23U83	Fromo sapiens hypothetical protein FLJ21128 (FLJ21128), IliRNA	e-121
1		Homo sapiens, transmembrane 4 superfamily member (tetraspan NET-	
1861	BC004161	7), clone MGC:2447 IMAGE:2958221, mRNA, complete cds	0.051
1001	B0004101	Homo sapiens, Similar to nucleolin, clone MGC:16354	0.031
1862	BC009855	IMAGE:3926227, mRNA, complete cds	e-171
1002	2000,000	Mus musculus adult male testis cDNA, RIKEN full-length enriched	0 1/1
1863	AK016507	library, clone:4931439C15, full insert sequence	0.019
1864	AF334756	Homo sapiens interleukin-1 HY2 (IL1HY2) gene, complete cds	1.5
		Human DNA sequence from cosmid L118G10, Huntington's Disease	
1865	Z68881	Region, chromosome 4p16.3	0.005
		Homo sapiens partial POLR2J2 gene for RPB11b1 protein, intron 4	
1866	AJ277737	(1824 BP)	0.0000001
1867	XM_051572	Homo sapiens heat shock transcription factor 2 (HSF2), mRNA	1E-11
1868	AF256918	Salmo salar clone BHMS267 microsatellite sequence	0.11
1869	AE000557	Helicobacter pylori 26695 section 35 of 134 of the complete genome	0.17
1870	XM_050469	Homo sapiens ferritin, light polypeptide (FTL), mRNA	0.081
1871	XM_042818	Homo sapiens homeo box D4 (HOXD4), mRNA	0.24
1872	17005000	Petunia x hybrida mRNA for lateral shoot inducing factor, complete	
18/2	AB035093	cds	1.9
1873	AE006184	Pasteurella multocida PM70 section 151 of 204 of the complete genome	5.6
10/3	AL000164	genome	3.0
1874	AE006880	Sulfolobus solfataricus section 239 of 272 of the complete genome	3.3
1875	NC_002669	Bacteriophage bIL310, complete genome	4.7
		Mus musculus 10, 11 days embryo cDNA. RIKEN full-length	- · · ·
1876	AK012924	enriched library, clone:2810048H02, full insert sequence	1.8
1877	XM_032188	Homo sapiens cell division cycle 27 (CDC27), mRNA	5.5
1878	M33662	Hamster G-o protein alpha subunit 2 mRNA, complete cds	5.8
1879	Y00478	L.esculentum LEACO2 gene	0.28
1880	XM_050519	Homo sapiens annexin A2 (ANXA2), mRNA	8E-34
1881	AF054822	Drosophila melanogaster activin beta precursor, gene, partial cds	0.43

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
1907	XM_012289	Homo sapiens cytoskeleton associated protein 2 (CKAP2), mRNA	2
1908	U44391	Human Down Syndrome region of chromosome 21, clone A12H1-1B2	0.008
1909	L04961	Mouse nuclear-localized inactive X-specific transcript (Xist) mRNA	1.8
1910	X61202	Mcthanococcus voltae fruA, fruB, fruG genes for subunits of F420- reducing hydrogenase and fruD orf	0.078
	1700000	Homo sapiens genomic DNA, chromosome 21q22.1, clone:f43D11,	
1911	AP000242	SOD-AML region, complete sequence	0.0001
1912	AL023777	S.pombe chromosome III cosmid c1827	0,66
1913	XM 038450	Homo sapiens hypothetical protein FLJ20694 (FLJ20694), mRNA	0
1914	AF129510	Mus musculus DACH protein (Dach) mRNA, complete cds	2.1
272.	12.27510	Plasmodium falciparum chromosome 2, section 42 of 73 of the	2.1
1915	AE001405	complete sequence	0.18
1916	XM_039566	Homo sapiens KIAA0948 protein (KIAA0948), mRNA	6.8
		Homo sapiens genomic DNA, chromosome 21q22.2, clone:T1492,	
1917	AP001431	LB7T-ERG region, complete sequence	0.074
1918	AF329137	Camelus bactrianus microsatellite Cms1 sequence	0.008
1919	NM_003816	Homo sapiens a disintegrin and metalloproteinase domain 9 (meltrin gamma) (ADAM9), mRNA	
1919	NN1_003816	Plasmodium falciparum chromosome 2, section 51 of 73 of the	1E-17
1920	AE001414	complete sequence	0.001
1111		Caenorhabditis elegans protein disulphide isomerase isoform I (pdi-1)	0.001
1921	U95074	and cyclophilin isoform 9 (cyp-9) genes, complete cds	0.029
1922	AB019534	Homo sapiens gene for cathepsin L2, complete cds	0.07
1923	AJ133777	Arabidopsis thaliana mRNA for gamma-adaptin 2	6.6
		Homo sapiens cyclin-dependent kinase inhibitor 2C (p18, inhibits	
1924	XM_001304	CDK4) (CDKN2C), mRNA	0.25
		Clostridium botulinum genes for ORF-22, HA-70, HA-17, HA-33,	
1925	AB037166	NTNHA, neurotoxin, complete cds	0.2
1926	Z48243	A.thaliana PARP mRNA for PARP protein	0.081
1927	XM_041006	Homo sapiens choreoacanthocytosis gene; KIAA0986 protein (KIAA0986), mRNA	0,25
		Homo sapiens uncharacterized hematopoietic stem/progenitor cells	
1928	XM_050592	protein MDS032 (MDS032), mRNA	0.083
1929	U34610	Mus musculus alpha-1(XVIII) collagen (COL18A1) gene, exon 6 and 7	0.008
		Homo sapiens, clone MGC:747 IMAGE:3343994, mRNA, complete	
1930	BC007658	cds	6E-50
		Human DNA sequence from clone RP11-45G20 on chromosome 6,	
1931	AL451135	complete sequence [Homo sapiens]	2
1932	AB004883	Marchantia paleacea mRNA for ribulose 1,5-bisphosphate	0.26
1932	AD004883	carboxylase/oxygenase small subunit, complete cds	0.26

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
1933	AF019973	Rattus norvegicus neuron-specific enolase (NSE) mRNA, complete cds	0.22
1934	AF231684	Drosophila melanogaster outstretched (os) gene, complete cds	0.75
1935	AC079136	Homo sapiens clone RP11-153L12, complete sequence	0.75
1936	AE006611	Streptococcus pyogenes M1 GAS strain SF370, section 140 of 167 of the complete genome	1.5
1937	AE007418	Streptococcus pneumoniae section 101 of 194 of the complete genome	0.029
1938	L26949	Simian immunodeficiency virus (T5) surface envelope glycoprotein proviral gene, 5' end	0.027
1939	AF343914	Campylobacter jejuni NCTC 11828 LOS biosynthesis cluster, partial sequence	0.58
1940	Z69925	Human DNA sequence from clone LL22NC03-116A5 on chromosome 22 Contains GSSs, complete sequence [Homo sapiens]	6.2
1941	U96876	Homo sapiens insulin induced protein 1 (INSIG1) gene, complete cds	0.78
		Homo sapiens similar to protein kinase C binding protein 1 (H.	
1942	XM 030024	sapiens) (LOC90218), mRNA	0.009
1943	AF226993	Rattus norvegicus selective LIM binding factor mRNA, complete cds	0.7
1944	AF033037	Plasmodium falciparum merozoite surface protein 4, merozoite surface protein 5, merozoite surface protein 2, and adenylosuccinate lyase genes, complete cds	0.69
1945	AB053087	Human immunodeficiency virus type 1 pol gene, reverse transcriptase amino terminal part, partial cds, 99 JP-NH3-II plasma virus clone:NH3RT-V3-2-9	0.028
1946		Homo sapiens hypthetical protein PRO2389 (PRO2389), mRNA	0.0001
1947	XM_010334	Homo sapiens transducin (beta)-like 1 (TBL1), mRNA	2.2
1948	AK003943	Mus musculus 18 days embryo cDNA, RIKEN full-length enriched library, clone:1110028F11, full insert sequence	2E-10
1949	AF085600	Drosophila melanogaster inorganic pyrophosphatase NURF-38 (Nurf- 38) gene, complete cds	0.78
1050	1170701	Plasmodium berghei extrachromosomal plastid PB-1, ORF470 gene, partial eds, tRNA-Thr, large subunit ribosomal RNA, tRNA-Met, tRNA-Arg, tRNA-Val, tRNA-Arg, tRNA-Leu, tRNA-Asn, tRNA-Ala,	. = .
1950	U79731	and small subunit ribosomal RNA genes, complete sequences	0.72
1951	AK022877	Homo sapiens cDNA FLJ12815 fis, clone NT2RP2002546	0.003
1952	M62755	Potato 4-coumarateCoA ligase (St4C1-1) gene, complete cds	0.085
1953	AK017902	Mus musculus adult male thymus cDNA, RIKEN full-length enriched library, clone:5830406J20, full insert sequence	2.3
		Sorghum bicolor maturase (matK) gene, complete cds; chloroplast	
1954	AF164418	gene for chloroplast product	0.078
1955	X54567	Human MFD22 dinucleotide repeat DNA	3E-13

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID	A COPPORT		
NO	ACCESSN	DESCRIP	P VALUE
1956	AB056832	Macaca fascicularis brain cDNA clone:QfIA-14079, full insert sequence	0,00009
	1220002	Homo sapiens ubiquitously transcribed tetratricopeptide repeat gene,	0.00009
1957	XM 034132	Y chromosome (UTY), mRNA	7
		Homo sapiens complement component 2 (C2) gene allele b, exons 1	
1958	L09706	through 8	0.24
1959	XM_017231	Homo sapiens hypothetical protein FLJ23342 (FLJ23342), mRNA	8E-76
1960	X83989	P.palustris chloroplast rbcL gene, promoter region	0.003
1961	AK025090	Homo sapiens cDNA: FLJ21437 fis, clone COL04285	0
10.0		Mus musculus adult male hippocampus cDNA, RIKEN full-length	
1962	AK013805	enriched library, clone:2900079F10, full insert sequence	0.14
1963	AE006210	Pasteurella multocida PM70 section 177 of 204 of the complete genome	
1963	AF075080		0.042
1965	D42118	Homo sapiens full length insert cDNA YQ80D07	0.000001
1965	D42118	Tobacco gene for parAs protein, promoter region and partial cds	0.009
1966	AF269350	Staphylococcus epidermidis strain SR1 clone step.1002d12 genomic sequence	2.3
1700	AI 200330	Staphylococcus aureus cell division protein FtsZ (ftsZ) gene, partial	2.3
		cds; YlmD (ylmD), YlmE (ylmE), YlmF (ylmF), YlmG (ylmG), and	Ì
		YlmH (ylmH) genes, complete cds; and cell division protein DivIVA	
1967	AF068904	(divIVA) gene, partial cds	0.19
1707	711 000701	Human DNA sequence from clone RP11-117A20 on chromosome 6,	0.19
1968	AL589920	complete sequence [Homo sapiens]	0.003
1700	122507720	Mus musculus survival motor neuron (Smn) gene, promoter region	0,003
1969	AF027688	and partial cds	0.56
		Homo sapiens mRNA; cDNA DKFZp564P056 (from clone	
1970	AL049340	DKFZp564P056)	0.25
1971	AJ223385	Fowlpox virus strain HP-440 DNA, isolate FP9, 14.6 kb fragment	0.25
		Human mitochondrial DNA control region, Indonesian, sequence 1 of	
1972	U25354	60	3E-65
		Staphylococcus epidermidis strain SR1 clone step.1027g02 genomic	
1973	AF269868	sequence	0,083
		Drosophila melanogaster genomic scaffold 142000013385598,	
1974	AE003326	complete sequence	0.087
1975	XM_050217	Homo sapiens RNA binding protein (LOC84549), mRNA	2.1
		Plasmodium falciparum chromosome 2, section 39 of 73 of the	
1976	AE001402	complete sequence	0.026
1977	XM_001935	Homo sapiens forkhead box D2 (FOXD2), mRNA	2.1
1000	AT 50020 :	Human DNA sequence from clone RP11-349A16 on chromosome	
1978	AL590384	Xq22.3-24, complete sequence [Homo sapiens]	0.008
1979	AL117050	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen	
1979	XM 037529	deprivation Homo sapiens KIAA0410 gene product (KIAA0410), mRNA	2.2
1980	XM_037529 X56212		
1981	A30212	C.elegans gene for vitellogenin	1.8

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	Process	
1982	AF231684	DESCRIP	P VALUE
1902	AF231064	Drosophila melanogaster outstretched (os) gene, complete cds	0.7
1983	AB032264	Danio rerio mRNA for glycogen synthase kinase 3 alpha, complete cds	2.2
1984	AK022877	Homo sapiens cDNA FLJ12815 fis, clone NT2RP2002546	0.009
1985	U89140	Mus musculus aldose reductase gene, promoter region and exon 1	0.69
1986	M37036	Rat nucleolar proteins B23.1 and B23.2	0.086
		Homo sapiens amiloride-sensitive cation channel 2, neuronal	
1987	XM_035498	(ACCN2), mRNA	0.25
1988	U40944	Caenorhabditis elegans cosmid PDB1, complete sequence	0.025
1989	AF102762	Dasylagon n. sp. 'Mardulyn & Whitfield' 16S large subunit ribosomal RNA gene, partial sequence; mitochondrial gene for mitochondrial product	0.24
		Maoricicada iolanthe cytochrome oxidase subunit I (COI) gene,	
1990	AF247631	partial cds; mitochondrial gene for mitochondrial product	0.026
1991	Z81538	Caenorhabditis elegans cosmid F45H10, complete sequence	6.2
1992	Z72516	Caenorhabditis elegans cosmid T25G3, complete sequence	0.75
1993	AE002201	Chlamydophila pneumoniae AR39, section 32 of 94 of the complete genome	6.6
1994	NM_003816	Homo sapiens a disintegrin and metalloproteinase domain 9 (meltrin gamma) (ADAM9), mRNA	2E-17
1995	AL158132	Human DNA sequence from clone RP3-323A24 on chromosome 4. Contains GSSs and a putative CpG island, complete sequence [Homo sapiens]	0.074
1996	NM 023122	Mus musculus glycoprotein m6b (Gpm6b), mRNA	5.7
1997	AF014502	Glycine max seed coat peroxidase precursor (Ep) gene, complete cds	1.8
1998	AJ270226	Entodinium caudatum partial mRNA for putative glycosyltransferase, clone L48	2,1
1999	AF159913	Euplotes crassus transposon Tecl clone Tecl-2 orf 2 and orf 3 pseudogenes, complete sequence	0.011
2000	AF101309	Caenorhabditis elegans cosmid H24G06, complete sequence	2.2
2001	AB021240	Hypseleotris compressus mitochondrial gene for Cytochrome b, complete cds	0.022
2001		Homo sapiens 3 BAC PAC-56F11 (Roswell Park Cancer Institute	0.022
2002	AC078798	Human BAC Library) complete sequence	0.001
2003	AK021504	Homo sapiens cDNA FLJ11442 fis, clone HEMBA1001327	0.76
2004	AF314531	Chilli leaf curl virus A component, complete sequence	0.23
2005	XM_047794	Homo sapiens KIAA1641 protein (KIAA1641), mRNA	e-123
2006	AC079136	Homo sapiens clone RP11-153L12, complete sequence	0.65
2007	AE001422	Plasmodium falciparum chromosome 2, section 59 of 73 of the	0.001
2007	AE001422 AJ306692	complete sequence	0.001
2009	AJ306692	Drosophila virilis ORF1, rpL14 gene and ORF2	0.26
2010	AC084242	Arabidopsis thaliana chromosome 1 BAC T24P22 genomic sequence, complete sequence	0.72

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID	. congan		
NO	ACCESSN	DESCRIP	P VALUE
2011	Z16768	H. sapiens (D12S90) DNA segment containing (CA) repeat, clone AFM172xd8; single read	0.058
2012	AF311633	Eimeria nieschulzi ORF470 gene, partial cds; apicoplast gene for apicoplast product	1.9
2013	XM_012933	Homo sapiens hypothetical protein FLJ11656 (FLJ11656), mRNA	2.5
2014	AB035190	Homo sapiens RHD gene, intron 2, complete sequence	1.6
2015	NC_002135	Pleurotus ostreatus mitochondrial plasmid mlp1, complete sequence	0.22
		Elgaria paucicarinata NADH dehydrogenase subunit I (ND1) gene, partial cds; tRNA-Ile, tRNA-Gln, and tRNA-Met genes, complete	
		sequence; NADH dehydrogenase subunit II (ND2) gene, complete cds:	
		tRNA-Trp, tRNA-Ala, tRNA-Asn, tRNA-Cys, and tRNA-Tyr genes,	
2016	AF085619	c>	2.3
2017	D85424	Homo sapiens gene for alphaS1-casein, 5'flanking region	0,077
2018	AE001138	Borrelia burgdorferi (section 24 of 70) of the complete genome	0.026
2019	BC007114	Homo sapiens, clone IMAGE:4295422, mRNA	0.23
2020	AF080510	Homo sapiens mannose-binding protein gene, exon 4 and complete	0.00001
2020	AF080310	Homo sapiens hypothetical protein DKFZp761J139 (DKFZp761J139),	0.00001
2022	XM_027941	mRNA .	0.003
2023	Z68996	H.sapicns mRNA for immunoglobulin kappa light chain VJ region (ID POM021)	0.15
		Mus musculus DNA, clone:lambda4/6, endogenous mouse mammary	
2024	AB049191	tumor virus LTR region	0.15
١ ١		No. of the state o	
		Mycoplasma capricolum NADH oxidase (naox) gene, partial cds, and lipoate-protein ligase (lpla), pyruvate dehydrogenase EI alpha subunit	
		(odpa), pyruvate dehydrogenase EI beta subunit (odpb), pyruvate	
2025	U62057	dchydrogenase EII (odp2), dihydrolipoamide dchydroge>	3
2023	002037	Oryza sativa microsatellite MRG0852 containing (AT)X17, genomic	
2026	AY018527	sequence	0.0003
		Phyllopetalia apicalis large subunit ribosomal RNA gene, partial	
1 1		sequence; tRNA-Valine gene, complete sequence; and small subunit	
-		ribosomal RNA gene, partial sequence; mitochondrial genes for	
2027	AF266084	mitochondrial products	0.009
		Cucumis melo galactinol synthase (GAS1) gene, upstream sequence	
2028	AF249912	and partial cds	0.021
		Homo sapiens hypothetical protein DKFZp434F1017	
2029	XM_048172	(DKFZP434F1017), mRNA	6.5
2030	ATO CTASS	Erysimum cheiri chloroplast outer envelope membrane protein	
2030	AF367436	mRNA, complete cds Homo sapiens kallikrein-like protein 3 (KLK9) gene, complete cds.	1.2
2031	AF135026	alternatively spliced	0.76
2031	Z69782	A.thermophilum xylR gene, xynA gene and alfA gene	0.0004
2002	207702	A. the mophism Ayre gone, Ayre gene and an A gene	0.0004

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DEGENER	Ī
NU	ACCESSN	DESCRIP	P VALUE
2060	AE002123	Ureaplasma urealyticum section 24 of 59 of the complete genome	1.8
2061	Z68019	HIV-1 DNA V3 region (homosexual H1031, seroconversion sample 1987)	0.25
2062	XM_006487	Homo sapiens spectrin, beta, non-erythrocytic 2 (SPTBN2), mRNA	0.000003
2063	AF329716	Oncorhynchus mykiss clone OSU9 T cell receptor beta chain variable region gene, partial cds	0.0008
2064	AC024773	Caenorhabditis elegans cosmid Y40C7A, complete sequence	0.035
2065	XM_032748	Homo sapiens KIAA1634 protein (KIAA1634), mRNA	0.079
2066	M58526	Human alpha-5 collagen type IV (COL4A5) mRNA, 3' end	2.1
2068	X56697	B.thuringiensis gene for sigma factor 35	0.59
2069	D00768	Pig pituitary glycoprotein hormone alpha subunit gene, exons 3 and 4 and 3 flank	1.6
		Boophilus microplus 12S ribosomal RNA gene, partial sequence; tRNA-Ile, tRNA-Gln, and tRNA-Phe genes, complete sequence; and NADH dehydrogenase subunit 5 (ND5) gene, partial cds,	
2070	AF110621	mitochondrial genes for mitochondrial products	0.21
2071	AC024779	Caenorhabditis elegans cosmid Y43B11AL, complete sequence	0.24
2072	AK023589	Homo sapiens cDNA FLJ13527 fis, clone PLACE1006076	0
		W.suaveolens mitochondrial DNA intergenic region between ORF1	
2073	X77691	and 15S rRNA genes	0.24
2074	Z68749	Caenorhabditis elegans cosmid F56H11, complete sequence	1.4
		Solanum tuberosum glutamine synthetase GS2 (gln) mRNA, partial	
2075	AF302113	cds; nuclear gene for plastid product	1.4
		Trichaptum abietinum mitochondrial small subunit ribosomal RNA,	
2076	U27078	mitochondrial gene, partial sequence	0.64
2077	XM_012456	Homo sapiens golgin-67 (KIAA0855), mRNA	2.2
2078	AB032960	Homo sapiens mRNA for KIAA1134 protein, partial cds	1E-36
		Homo sapiens hypothetical gene supported by AF106046; AK000332	
2079	XM_052376	(LOC93587), mRNA	0.66
2080	AB021866	Homo sapiens KIP gene, complete cds	1.3
2081	XM_029213	Homo sapiens hypothetical protein DKFZp564O0523 (DKFZP564O0523), mRNA	0,38
2082	AF384143	Triticum aestivum pathogenesis-related protein 1 mRNA, complete cds	_0.0003
	12000000	Neisseria meningitidis serogroup B strain MC58 section 10 of 206 of	
2083	AE002368	the complete genome	0.71
2084	AE006662	Sulfolobus solfataricus section 21 of 272 of the complete genome	0.63
2085	AY020760	Oryza sativa microsatellite MRG3085 containing (TA)X13, closest to marker R2976, genomic sequence	6.9
2086	AF161419	Homo sapiens HSPC301 mRNA, partial cds	1.8
2087	AJ249838	Gallus gallus mRNA for radixin (rdx gene)	2.1
2007	AV2-17030	Corynebacterium glutamicum heat shock, ATP-binding protein (clpB)	2.1
2088	U43536	gene, complete cds	0.22
2089	XM_037588	Homo sapiens KIAA0874 protein (KIAA0874), mRNA	2E-34

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
2090	NC 002697	Chrysomya chloropyga mitochondrion, complete genome	0.077
2091	AF332207	Caenorhabditis elegans clone yk509b7 nuclear receptor NHR-57 gene, partial cds	0.45
2092	U39886	Bos taurus Y-chromosome specific genomic sequence	0.2
2093	Z19055	B.aphidicola tryptophan operon	0.26
2094	AJ290309	Simethis mattiazzii chloroplast trnL-trnF intergenic spacer	0.57
2095	AK013631	Mus musculus adult male hippocampus cDNA, RIKEN full-length enriched library, clone:2900041A09, full insert sequence	0.18
2096	AF015468	Plasmodium falciparum microsatellite LT103 sequence	0.11
2097	AJ277734	Sus scrofa microsatellite sequence S0601	0.4
2098	AF203341	Glycine max chloroplast carboxyl transferase alpha subunit (accA-4) nuclear pseudogene, partial sequence; and putative steroid reductase gene, complete cds	0,001
2099	XM_052597	Homo sapiens KIAA1350 protein (KIAA1350), mRNA	0
2100	Z96649	H.sapiens telomeric DNA sequence, clone 6QTEL010, read 6QTEL00010.seq	0.007
2101	BC009105	Mus musculus, Harvey rat sarcoma oncogene, subgroup R, clone MGC:6162 IMAGE:3483313, mRNA, complete cds	4.1
2102	AF245117	Mus musculus leukocyte cell-surface molecule (Ly9) gene, exon 1	0.025
2103	AL023810	Caenorhabditis elegans cosmid C16D2, complete sequence	0.008
2104	X17051	E.gracilis DNA for ribosomal protein operon	0.21
2105	AF005392	Homo sapiens alpha tubulin (TUBA2) gene, partial cds	0.63
2106	AE002798	Drosophila melanogaster genomic scaffold 142000013385466, complete sequence	0.025
2107	AE002114	Ureaplasma urealyticum section 15 of 59 of the complete genome	0.074
2108	L48039	Fibrobacter succinogenes endo-1,4-beta-D-glucanase gene, complete cds, endo-1,4-beta-D-glucanase gene, 5' end of cds	2
2109	D87922	Rat DNA for 3'UTR of skeletal muscle sodium channel, partial sequence	0.71
2110	AF157137	Gongronella butleri 18S ribosomal RNA gene, partial sequence	0.24
2111	Z92970	Caenorhabditis elegans cosmid H06O01, complete sequence	0.24
2112	AB035494	Alternaria alternata gene for Akt3-2, complete cds	0.61
2113	M38351	Rat embryonic glutamic acid decarboxylase gene, embryonic stop (ES) exon	0.67
2114	X16715	D. melanogaster gene for chorion protein s16	0.21
2115	U16271	Homo sapiens AMP deaminase isoform L (AMPD2) gene, exons 1A and 1B	0.07
2116	AF317672	Helicobacter pylori isolate 4424 vacuolating cytotoxin VacA (vacA) gene, partial cds	0.24
2117	NM_010250	Mus musculus gamma-aminobutyric acid (GABA-A) receptor, subunit alpha 1 (Gabra1), mRNA	0.76
2118	BC010786	Mus musculus, Similar to CREB/ATF family transcription factor, clone MGC: 18836 IMAGE:4211480, mRNA, complete cds	0.69

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ano m		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID	ACCESSN	Diagram	
NO	ACCESSIN	DESCRIP	P VALUE
2119	U67482	Methanococcus jannaschii section 24 of 150 of the complete genome	0.24
2120	U73373	HIV-1 clone 1063-2-VPR vpr gene, complete cds	2
2121	AE007370	Streptococcus pneumoniae section 53 of 194 of the complete genome	0.68
J l		Human DNA sequence from clone RP11-533E16 on chromosome 6,	
2122	AL449163	complete sequence [Homo sapiens]	2E-11
1 ,,,,, 1	37.00000	Antithamnion sp. rhodoplast genes atpl, atpH, atpG, atpF, atpD, atpA,	
2123	X63382 AF186797	orf1, orf2 and orf3	0.081
2124	AF186/9/	Homo sapiens RalGDS-like (RGL) gene, exon 17	0.009
2125	AF110796	Rattus norvegicus GABAB1 receptor (GABABR1) gene, exons 1 through 9	
2126	XM 038798	Homo sapiens KIAA1457 protein (KIAA1457), mRNA	4.2 2E-96
2127	U96998	Homo sapiens MET proto-oncogene, intron 19, 3' end	0.67
2127	0,0,,,,	Lactococcus lactis subsp. lactis IL1403 section 172 of 218 of the	0.67
2128	AE006410	complete genome	0.084
2129	AB050422	Macaca fascicularis brain cDNA, clone:QnpA-21421	6.9
	112030422	Mus musculus medium and short chain L-3-hydroxyacyl-Coenzyme A	0.5
		dehydrogenase (Mschad) gene, exons 2 through 8, and complete cds;	
2130	AF375597	nuclear gene for mitochondrial product	2.2
		Drosophila melanogaster genomic scaffold 142000013385931,	
2131	AE002758	complete sequence	0.26
2132	AJ008105	Chrysolina timarchoides COI gene, partial CDS	0.031
2133	U67526	Methanococcus jannaschii section 68 of 150 of the complete genome	2
2134	AJ223966	Mus musculus NAGA gene	0.63
2135	AF016986	Callicebus moloch gammal-globin gene, complete cds	0.68
2136	U66872	Danio rerio enhancer of rudimentary homolog mRNA, complete cds	0.000
2137	AJ271220	Homo sapiens partial gene for dystrophin, intron 49	0.029
2137	A32/1220	Canis familiaris prostaglandin E2 receptor subtype EP2 mRNA,	0.009
2138	AF075602	complete cds	0.23
2150	12 073002	Somplete cus	0.23
2139	XM_045015	Homo sapiens hypothetical protein FLJ00052 (FLJ00052), mRNA	0.75
2140	AY034614	Danio rerio Sec61 alpha form B mRNA, complete cds	0.1
		Homo sapiens similar to MACROPHAGE MANNOSE RECEPTOR	
2141	XM_028150	PRECURSOR (H. sapiens) (LOC90010), mRNA	1.9
2142	M24026	Rat MHC class I RT1 (RT44) mRNA (u haplotype), 3' end	0.24
		Mus musculus adult male tongue cDNA, RIKEN full-length enriched	
2143	AK010194	library, clone:2310076E16, full insert sequence	2
2144	AF165062	Hepatitis C virus strain MD9-2 complete genome	1.2
2145	AE007469	Streptococcus pneumoniae section 152 of 194 of the complete genome	2.1
1 2146	707240	Plasmodium falciparum DNA *** SEQUENCING IN PROGRESS	
2146 2147	Z97349 NM 010161	*** from contig 3-06, complete sequence	0.046
214/	TATAT_OTOTO1	Mus musculus ecotropic viral integration site 2 (Evi2), mRNA	0.02

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WC0214560 [fle://E/WO0214500 cpc]

Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE Escherichia coli O157:H7 EDL933 genome, contig 3 of 3, section 205 AE005586 2148 of 290 1.2 Psacothea hilaris isolate HA15 16S ribosomal RNA gene, partial 2149 AF332928 sequence; mitochondrial gene for mitochondrial product 0.069 2150 D49504 Borrelia garinii gene for outer surface protein C, complete cds Homo sapiens fibroblast growth factor receptor gene (located in the 2151 L25647 central MHC) signal peptide and consecutive exon 0.05 2152 AB040746 Carassius auratus gene for c-MYC, complete cds 0.063 X55978 T. brucei ESAG 8 gene for a leucine-rich repeat family protein 2154 AB010426 Phytoplasma sp. gene for AL1 like protein, complete cds 0.009 2155 XM 005841 Homo sapiens cdk inhibitor p21 binding protein (TOK-1), mRNA 5.8 2156 AK025222 Homo sapiens cDNA: FLJ21569 fis. clone COL06508 1E-33 Mesembryanthemum crystallinum protein kinase MK5 mRNA, 2157 AF234652 complete cds 0.2 Streptococcus pyogenes M1 GAS strain SF370, section 42 of 167 of 2158 AE006513 the complete genome 9.2 2159 XM 003660 Homo sapiens calcium modulating ligand (CAMLG), mRNA 0.023 2160 AF003490 Mesostoa kerri 16S ribosomal RNA gene, partial sequence 0.000004 2161 XM_048768 Homo sapiens tetratricopeptide repeat domain 3 (TTC3), mRNA 6.7 Plasmodium yoelii yoelii clone 10 235 kDa rhoptry protein (IIIa.1) 2162 AF323442 gene, partial cds 0.019 Mus musculus adult male testis cDNA, RIKEN full-length enriched 2163 AK016560 library, clone:4932441H21, full insert sequence 47 2164 Y12332 E.tenella plastid genes rps12, rps7, tuf, ORF45, and tRNA-Phe 0.003 Botrytis cinerea strain T4 cDNA library under conditions of nitrogen 2165 AL114081 deprivation 0.002 Caenorhabditis elegans clone yk509b7 nuclear receptor NHR-57 gene, 2166 AF332207 0.49 Helicoverpa zea clone S211 cytochrome P450 (CYP6B8) mRNA 2167 AF102263 complete cds 0.074 Human DNA sequence from clone RP1-213J1p on chromosome 2168 AL133397 22q11.21-12.2, complete sequence [Homo sapiens] 0.039 Homo sapiens mRNA: cDNA DKFZp761O0511 (from clone 2169 AL157423 DKFZp761O0511) 7E-15 2170 AF282578 Homo sapiens clone 20ptel c1622 12 3.3f sequence 9E-17 Oryzias latipes gene for soluble guanylyl cyclase alpha subunit, 2171 AB022280 complete cds 3.7 Homo sapiens transcriptional repressor p54 gene, promoter region and 2172 AF052744 2 2173 XM 007685 Homo sapiens MAGE-like 2 (MAGEL2), mRNA 0.008 Mus musculus adult male testis cDNA, RIKEN full-length enriched 2174 AK019539 library, clone:4921514G21, full insert sequence 1.6 2175 AE007472 Streptococcus pneumoniae section 155 of 194 of the complete genome 6.2 Human DNA sequence from clone RP11-174B19 on chromosome 6. 2176 AL162381 complete sequence [Homo sapiens] 0.059

WO 02/14500 PCT/US01/25840

		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Mouse extra-embryonic endodermal cytokeratin type II (EndoA)	
2177	M21836	mRNA, complete cds	e-138
2178	X06292	Human c-fes/fps proto-oncogene	1.7
		Arabidopsis thaliana putative MLH1 protein (AT4g09140) mRNA,	
2179	AF360278	complete cds	0.0001
2180	X78548	G.max gene encoding epoxide hydrolase	0.68
		H.sapiens CpG island DNA genomic Mse1 fragment, clone 97b6,	
2181	Z56222	reverse read cpg97b6.rt1a	5E-60
		Wiebesia brusi cytochrome oxidase subunit I (COI) gene, partial cds;	
- 1		tRNA-Leu gene, complete sequence; and cytochrome oxidase subunit	
- 1		II (COII) gene gene, partial cds; mitochondrial genes for	
2182	AF200412	mitochondrial products	0.084
		Arabidopsis thaliana putative metal-binding protein	
2183	AF360327	(MLE2.16/AT5g63530) mRNA, complete cds	0.014
		Homo sapiens CUG triplet repeat, RNA-binding protein 2 (CUGBP2).	
2184	XM_005837	mRNA	0.22
		Mus musculus, caspase 1, clone MGC:6106 IMAGE:3583883,	
2185	BC008152	mRNA, complete cds	1.9
		Dictyostelium discoideum K7 kinesin-like protein mRNA, complete	
2186	U41289	cds	0.66
2187	AF163863	Mustela vison tyrosine aminotransferase gene, complete cds	0.23
\neg		, , , , , , , , , , , , , , , , , , , ,	
2188	XM_017198	Homo sapiens hypothetical protein FLJ12085 (FLJ12085), mRNA	4E-36
2189	D16180	Human PMP2 gene for peripheral myelin protein 2, exon 2 and 3	0.68
		Clostridium botulinum genes for ORF-22, HA-70, HA-17, HA-33,	
2190	AB037920	NTNHA, neurotoxin, complete cds	0.083
2191	AE006703	Sulfolobus solfataricus section 62 of 272 of the complete genome	1.6
		The state of the s	
2192	AB047401	Periplaneta americana Vg-2 mRNA for vitellogenin-2, complete cds	0.25
		Human rotavirus M37 non-structural protein NSP1 mRNA, complete	0.20
2193	U11491	cds	0.17
2194	AF338818	Mus musculus flavohemoprotein b5/b5R mRNA, complete cds	2.3
		Mus musculus genomic DNA, chromosome 7, clone:B118 5,	2.5
2195	AP001290	complete sequence	0.56
2196	XM 010940	Homo sapiens putative GR6 protein (GR6), mRNA	2E-61
		Plasmodium falciparum para-aminobenzoic acid synthetase gene.	ZLI VI
2197	AF119554	complete cds	0,009
		Streptococcus pneumoniae strain SPN1506 topoisomerase IV subunit	0.002
2198	AY035995	A (parC) gene, complete cds	1.2
		Homo sapiens cAMP responsive element binding protein-like 2	1.2
2199	XM_006892	(CREBL2), mRNA	0.049
		Borrelia burgdorferi repeated DNA element, 30.5 kb circular plasmid	0,049
2200	X87127	copy	0.46
-200	280/12/	Clostridium thermocellum xynA and xynB genes for xylanase,	0.40

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
2202	AE003004	Drosophila melanogaster genomic scaffold 142000013385492, complete sequence	0.23
2203	U63418	Mus musculus uncoupling protein (Ucp) gene, nuclear gene encoding mitochondrial protein, complete cds	0.23
2204	AJ243265	Homo sapiens partial PGM1 gene for phosphoglucomutase 1, exons 5-7	e-135
2205	X13883	Yersinia pseudotuberculosis virulence plasmid plBI yopA gene for Yop1 protein	2.1
2206	XM 011639	Homo sapiens KIAA0535 gene product (KIAA0535), mRNA	8E-22
2207	AF098990	Caenorhabditis elegans cosmid H35N03	2.2
2208	AB037920	Clostridium botulinum genes for ORF-22, HA-70, HA-17, HA-33, NTNHA, neurotoxin, complete cds	0.2
2209	AB053087	Human immunodeficiency virus type 1 pol gene, reverse transcriptase amino terminal part, partial cds, 99JP-NH3-II plasma virus clone: NH3RT-V3-2-9	0.028
2210	NM_008989	Mus musculus purine rich element binding protein A (Pura), mRNA	0.028
2211	U97291	Equus caballus random genomic clone JH226	0.008
2212	AK001699	Homo sapiens cDNA FLJ10837 fis, clone NT2RP4001260, highly similar to Homo sapiens mRNA for KIAA0875 protein	0.016
2213	U84532	Human dystrobrevin (DTN) gene, exon 4	0.23
2214	Z54522	H.sapiens CpG island DNA genomic Mse1 fragment, clone 12b6, forward read cpg12b6.ft1c	1.5
		Human DNA sequence from cosmid 24F8 from a contig from the tip of the short arm of chromosome 16, spanning 2Mb of 16p13.3.	
2215	Z69666	Contains ESTs, repeat polymorphism and CpG island	6.5
2216	M69205	Ockelbo virus complete genome	2,2
2217	AF202552	Homo sapiens DNA methyltransferase (DNMT1) gene, exons 2, 3, and 4	0.24
2218	D16541	Chicken DNA for connectin(titin), partial sequence	0.064
2220	AC023351	Homo sapiens clone RP11-637J20, complete sequence	1E-59
2221	AL512305	Human DNA sequence from clone RP11-85E24 on chromosome 6, complete sequence [Homo sapiens]	0,002
2222	U67581	Methanococcus jannaschii section 123 of 150 of the complete genome	5
2223	AE007428	Streptococcus pneumoniae section 111 of 194 of the complete genome	1.9
2224	AK023601	Homo sapiens cDNA FLJ13539 fis, clone PLACE1006640	1.7
2225	AF375468	Homo sapiens endothelial protein C receptor (PROCR) gene, complete cds	0.016
2226	AF350726	Symphyglossum sanguineum chloroplast tRNA-Leu (trnL) gene and trnL-trnF intergenic spacer, partial sequence	1.4
2227	NM_008308	Mus musculus 5-hydroxytryptamine (scrotonin) receptor 1A (Hirla), mRNA	0.23
2228	Z62421	H. sapiens CpG island DNA genomic Mse1 fragment, clone 68h2, forward read cpg68h2.ft1a	4E-68

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WC0214500 [flig //E /WO0214500 opc]

		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		A TOMBON (DIEDLE 1 TO COMBINE)	
NO	ACCESSN	DESCRIP	P VALUE
2229	AF210834	Homo sapiens HARP (HARP) gene, exons 3 and 4	0.008
		Homo sapiens (subclone 2 b12 from P1 H49) DNA sequence,	
2230	L81672	complete sequence	0.13
		Homo sapiens integrin, alpha 3 (antigen CD49C, alpha 3 subunit of	
2231	XM_008431	VLA-3 receptor) (ITGA3), mRNA	1.8
		Lactococcus lactis subsp. lactis IL1403 section 57 of 218 of the	
2232	AE006295	complete genome	0.19
2233	U39693	Mycoplasma genitalium section 15 of 51 of the complete genome	0.21
2234	AL032629	Caenorhabditis elegans cosmid Y102F5A, complete sequence	0.46
		Ascogaster sp. 16S ribosomal RNA gene, mitochondrial gene for	
2235	AF029114	mitochondrial RNA, partial sequence	0.076
		Hyalomma dromedarii 12S small ribosomal RNA gene, partial	
2236	AF150036	sequence; mitochondrial gene for mitochondrial product	0.63
		Homo sapiens 22 kDa actin-binding protein (SM22) gene, complete	
2237	AF013711	cds	0,6
2238	XM_035961	Homo sapiens hypothetical protein FLJ10849 (FLJ10849), mRNA	0.77
		Drosophila erecta partial eIF2g gene for eukaryotic translation	
		initiation factor 2 gamma and Su(var)3-9 gene for heterochromatin	
2239	AJ290957	protein	0.24
2240	AF282004	Homo sapiens clone 15qtel_c366at7 sequence	8.2
2241	ATT104015	Plasmodium falciparum protein serine/threonine kinase-1 (psk1) gene,	0.000
2241	AF104915	Complete cds Mouse MHC class I Lyt-2-a gene encoding lyt-2.1 T-cell surface	0.009
2242	M22064	alloantigen, complete cds	0.049
2242	1912/2004	Hypseleotris compressa 12S ribosomal RNA gene, complete sequence;	0.049
2243	AF265368	mitochondrial gene for mitochondrial product	0.21
2243	AF203306	Plasmodium falciparum chromosome 2, section 34 of 73 of the	0.21
2245	AE001397	complete sequence	0.024
2210	122001037	Mus musculus adult male lung cDNA, RIKEN full-length enriched	0.021
2246	AK004676	library, clone:1200009I24, full insert sequence	0.56
		Homo sapiens sterile-alpha motif and leucine zipper containing kinase	- 1111
2247	XM 002436	AZK (ZAK), mRNA	0,67
		Lactococcus lactis subsp. lactis IL1403 section 86 of 218 of the	
2248	AE006324	complete genome	0.47
2249	XM 042833	Homo sapiens KIAA0295 protein (KIAA0295), mRNA	5.1
2250	X63412	B.burgdorferi plasmid ospA gene for outer surface protein A	0.36
2251	AK024235	Homo sapiens cDNA FLJ14173 fis, clone NT2RP2002755	1.8
		Eryx miliaris nogajorum cytochrome b (cytb) gene, mitochondrial	
2252	U69826	gene encoding mitochondrial protein, partial cds	5.9
		HIV-1 isolate TZB0036 from Tanzania, gp120 C2-C4 region (env)	
2254	AF038086	gene, partial cds	0.077
		Adoncholaimus thalassophygas isolate Sou39 large subunit ribosomal	
		RNA gene, partial sequence; mitochondrial gene for mitochondrial	
2255	AF317082	product	0.003
2256	NM_005386	Homo sapiens neuronatin (NNAT), mRNA	0.6

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WC0214560 [fle://E/WO0214500 opc]

Table 3A Nearest Neighbor (BlastN vs. Genbank) SEQ ID NO ACCESSN DESCRIP P VALUE 2257 XM 037301 Homo sapiens hypothetical protein FLJ12697 (FLJ12697), mRNA 2E-61 2258 XM 028046 Homo sapiens KIAA0220 protein (KIAA0220), mRNA 5E-61 Lactococcus lactis plasmid pND861 abortive phage resistance proteins 2259 U94520 (abiLi) and (abiLii) genes, complete cds 0.81 2260 1100037 Caenorhabditis elegans cosmid T20H4 0.081 2261 Z74866 S.cerevisiae chromosome XV reading frame ORF YOL124c 0.55 2262 Z92779 Caenorhabditis elegans cosmid C44E1, complete sequence 0.7 Saccharomyces cerevisiae mitochondrion COX/OXI3 gene encoding 2263 V00694 for cytochrome oxidase subunit 1 0.021 2264 AC010728 Homo sapiens BAC clone RP11-258E22 from Y, complete sequence 3E-30 2265 AJ406392 Beta vulgaris partial En/Spm-like transposon, clone 44.9 Plasmodium falciparum chromosome 2, section 67 of 73 of the 2266 AE001430 complete sequence 0.064 AF378270 2267 Pangasius sp. ZZSH-2001 microsatellite PSP-G 456 sequence Homo sapiens hypothetical gene supported by AL122112 2268 XM 047227 (LOC92776), mRNA 0.38 Drosophila melanogaster genomic scaffold 142000013385931. 2269 AE002758 complete sequence 0.2 Botrytis cinerea strain T4 cDNA library under conditions of nitrogen 2270 AL115861 deprivation 0.14 Lactococcus lactis subsp. lactis IL1403 section 169 of 218 of the 2271 AE006407 complete genome 1 8 2272 XM 049351 Homo sapiens KIAA1600 protein (KIAA1600), mRNA 0.0007 2273 AE005977 Caulobacter crescentus section 303 of 359 of the complete genome 5.2 Homo sapiens chromosome 12 open reading frame 5 (C12orf5), 2274 XM 007057 0.63 Homo sapiens hypothetical gene supported by AK023761 2275 XM 031923 (LOC90466), mRNA 0.0000007 2276 AJ005330 pGAII(-) SK positive selection cloning vector gltS gene 3E-21 2277 XM 029246 Homo sapiens collagen, type I, alpha 2 (COL1A2), mRNA 0.051 Homo sapiens polymerase delta small subunit (POLD2) gene, partial 2278 AF185279 scauence 0.002 2279 AF080592 Mus musculus centrin (Cetn2) gene, complete cds 1.8 2280 XM 033626 Homo sapiens nucleoporin-like protein 1 (NLP 1), mRNA 3.6 Ipomoea purpurea genes for dihydroflavonol 4-reductase, complete 2281 AB011667 0.25 Ipomoca purpurea genes for dihydroflavonol 4-reductase, complete 2282 AB011667 0.25 2283 XM 002698 Homo sapiens LOC89635 (LOC89635), mRNA 6.3 2284 BC009385 Homo sapiens, clone IMAGE:4127835, mRNA 0.61 2285 U39718 Mycoplasma genitalium section 40 of 51 of the complete genome 3,4 Schizosaccharomyces pombe spo14/stl1 gene for Sec12-like protein, 2286 AB036755 0.079 complete cds

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Sepiella maindroni isolate c 16S ribosomal RNA gene, partial	
2287	AF369960	sequence; mitochondrial gene for mitochondrial product	0.22
		Escherichia coli O157:H7 EDL933 genome, contig 2 of 3, section 29	l
2288	AE005360	of 55	0.6
		Eastern equine encephalomyelitis virus North American antigenic	
	******	variety nonstructural polyprotein and structural polyprotein genes,	_
2289	U01034	complete cds	1.7
2290	4 To 55100	Pleurotus ostreatus linear mitochondrial plasmid mlp2, partial	
2290	AF355103	sequence	5.7
2201	AC000043	Homo sapiens Chromosome 22q13 Cosmid Clone p74a8, complete	
2291	X89398	sequence	0.085
2292	A89396	H.sapiens ung gene for uracil DNA-glycosylase	0.71
2293	AY008264	Yersinia enterocolitica phage shock protein locus, complete sequence	2.2
2293	A 1 008204	Drosophila melanogaster Su(P) and anon-73B1 genes and partial o25	2.2
2294	AJ011320	gene and Pros26 gene	6,5
2234	AJ011320	Homo sapiens glutamate receptor, ionotropic, AMPA 1 (GRIA1),	0.3
2295	XM 040290	mRNA	4
2296	AJ293578	Homo sapiens partial MOCS1 gene, exons 2-8	6.3
22,0	113273376	Homo sapiens, clone MGC:12465 IMAGE;3683051, mRNA,	0.5
2297	BC007051	complete cds	7E-32
2298	M23103	B.subtilis spoIVCB gene, complete cds	0.2
2299	AF280606	Triticum aestivum omega gliadin pseudogene, complete sequence	0.24
		Platycnemis pennipes large subunit ribosomal RNA gene, partial	
]		sequence; tRNA-Valine gene, complete sequence; and small subunit	
		ribosomal RNA gene, partial sequence; mitochondrial genes for	
2300	AF266095	mitochondrial products	0.077
l .		Saccharomyces pastorianus CBS1538 small subunit ribosomal RNA	
2301	AF114927	gene, mitochondrial gene for mitochondrial RNA, complete sequence	0.23
2302	M68901	P.multocida adenylate cyclase (cya) gene, complete cds	0.23
2303	AJ235878	Amorphophallus kiusiuensis chloroplast atpB-rbcL spacer, isolate PL1	0.23
		Homo sapiens erythrocyte transmembrane protein (LOC51145),	
2304	NM_016158	mRNA	5E-34
2305	Y17254	Euplotes octocarinatus gamma-tubulin 2 micronuclear gene	2.3
2306	X99872	Sulfolobus acidocaldarius pyrB gene	0.033
2307	X81001	H.sapiens HCG II mRNA	0.001
2308	Z80213	Caenorhabditis elegans cosmid C09E9, complete sequence	0.025
2309	AF257022	Salmo salar clone BHMS7-009 microsatellite sequence	0.74
		W PNI C PDI TOOTIO	
2310	AL138748	Human DNA sequence from clone RP4-732E19 on chromosome	0.21
2310	AL 138/48	Xq21.1-21.33 Contains GSSs, complete sequence [Homo sapiens]	0.21
2311	AJ409108	Entamoeba histolytica rpl27a-2 gene for ribosomal protein large subunit 27a-2, exons 1-2	0.009
2312	U26556	Human ferritin H (FTHL13) pseudogene	0.009
2312	020550	Imman remain it (F FFIL 15) pseudogene	0.008

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			I
NO	ACCESSN	DESCRIP	P VALUE
		Protoclythia modesta 12S ribosomal RNA gene, partial sequence;	
2313	AF126320	mitochondrial gene for mitochondrial product	0.26
		Homo sapiens mRNA; cDNA DKFZp434K0610 (from clone	
2314	AL137590	DKFZp434K0610)	2
2315	Z81471	Caenorhabditis elegans cosmid C14B4, complete sequence	0.029
		Helicobacter pylori, strain J99 section 128 of 132 of the complete	
2316	AE001567	genome	0.028
		Tribolium freemani Woot retrotransposon, partial sequence and 3'	
2317	U47913	insertion junction F3(1)	0.081
	**********	Homo sapiens centrosomal P4.1-associated protein; uncharacterized	
2318	XM_037521	bone marrow protein BM032 (BM032), mRNA	2.2
2210	A F2 C0 F0 1	Staphylococcus epidermidis strain SR1 clone step.1017g02 genomic	
2319	AF269701	sequence	0.7
2320	AF378183	Oryza sativa cultivar Milyang23 cytosolic fructose-1 mRNA, partial cds	
2320	AF3/0103	Oryza sativa microsatellite MRG1415 containing (AT)X26, genomic	6.9
2321	AY019090	oryza sauva microsatemie MRG1415 containing (A1)X26, genomic secuence	0.085
2321	A1019090	Human DNA sequence from clone RP11-533E16 on chromosome 6,	0.085
2322	AL449163	complete sequence [Homo sapiens]	0.00004
LJLL	ALAHYIOS	complete sequence [1101110 saprens]	0.00004
		Adenovirus type 41 DNA-binding protein (DBP) gene, exons 2 and 3.	
2323	M19540	23K protease, complete cds, hexon and 100K protein genes, last exons	0.026
2020	2,22,5 10	Homo sapiens partial CD30 gene for cytokine receptor CD30 and	0.020
2324	AJ272029	promoter region	0.003
2325	AF082966	Homo sapiens genomic sequence	0.75
		Drosophila melanogaster genomic scaffold 142000013385820,	
2326	AE002940	complete sequence	0.028
		Human DNA sequence from cosmid L129H7, Huntington's Disease	
2327	Z54147	Region, chromosome 4p16.3 contains CpG island	0.001
2328	D12590	Rattus norvegicus DNA, RFLP marker at locus A403	0.077
2329	XM_035524	Homo sapiens KIAA1201 protein (KIAA1201), mRNA	0.025
		Homo sapiens soluble liver antigen/liver pancreas antigen	
2330	XM_003575	(LOC51091), mRNA	4E-27
		Homo sapiens, G1 to S phase transition 1, clone MGC:1735	
2331	BC009503	IMAGE:2822947, mRNA, complete cds	0.69
		Mus musculus, golgi vesicular membrane trafficking protein p18,	
2332	BC005572	clone MGC:11601 IMAGE:3967968, mRNA, complete cds	0.78
2333	U39726	Mycoplasma genitalium section 48 of 51 of the complete genome	0,086
2334	XM_039942	Homo sapiens Meis (mouse) homolog 3 (MEIS3), mRNA	0.72
2335	X67506	C. thermocellum ancA gene	0.084
2336	AT 117552	Homo sapiens mRNA; cDNA DKFZp564F1171 (from clone	0.22
2336	AL117553	DKFZp564F1171)	0.22
2337	AF115574	Pisum sativum pathogenesis-related protein (DRR206-c) gene, complete cds	0.009
2338	Y11874	M.musculus uPA gene, promoter sequence	2.3
2330	1110/4	INLIMUSCUMUS MEA GENE, PROMOTER SEQUENCE	

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
2339	BC008277	Mus musculus, Similar to Endothelin receptor type A, clone MGC:6567 IMAGE:2812426, mRNA, complete cds	2.2
2340	XM_009543	Homo sapiens transcription factor AP-2 gamma (activating enhancer- binding protein 2 gamma) (TFAP2C), mRNA	0,21
2341	NM_015809	Mus musculus keratin-associated protein 5-4 (Krtap5-4), mRNA	0.42
2342	AE000813	Methanobacterium thermoautotrophicum from bases 206991 to 217929 (section 19 of 148) of the complete genome	5.2
2343	XM_034632	Homo sapiens hypothetical protein PRO2832 (PRO2832), mRNA	0.0000005
2344	XM_034229	Homo sapiens similar to hypothetical protein FLJ14260 (H. sapiens) (LOC90803), mRNA	0.069
2345	XM_033361	Homo sapiens 62537 (FLJ00024), mRNA	0.008
2346	BC005332	Homo sapiens, Similar to immunoglobulin kappa constant, clone MGC:12418 IMAGE:3934658, mRNA, complete cds	3E-33
2347	AF321120	Mus musculus gap junction membrane channel protein alpha 8 (Gja8) gene, complete cds	2.1
2348	AB054512	Berardius bairdii DNA, SINE flanking sequence Tuti35 locus	0.77
2349	Z98549	Plasmodium falciparum DNA *** SEQUENCING IN PROGRESS *** from contig 3-14, complete sequence	0.029
2351	XM_033483	Homo sapiens hypothetical gene supported by L39957; L43086; M82526, M9093. S6182; S7391; L107999; L19763; U96290; U96294; U96397; X58082; Z18327; AB006847; AB027436; AB027438, AF007572; AF035035; AF0252335; AF099196; AF103384; AF103309; AF103466; AF10>	8E-33
2352	XM_033483	Homo sapiens hypothetical gene supported by L39957; L43086; M83256; M99603; S56182; S73911; U07989; U79763; U96290; U36294; U36397; X36082; Z18327; AB006847; AB027436; AB027438; AF007572; AF035035; AF035235; AF099196; AF103384; AF103390; AF103466; AF10>	1E-27
2353	XM 033483	Homo supiens lypothetical gene supported by L39957; L4,3086; M85256; M99603; S56182; S73911; U07989; U79763; U96290; U36294; U96397; X58082; Z18327; AB006847; AB027436; AB027438; AF007572; AF035035; AF0252353; AF099196; AF103384; AF103394; AF103394; AF10466; AF10>	av. 22
2555	AM_033463	Homo sapiens hypothetical gene supported by AF074988	8E-33
2354	XM_043728	(LOC92227), mRNA	0.001
2355	AE001425	Plasmodium falciparum chromosome 2, section 62 of 73 of the complete sequence	6.4
2356	XM_033483	Homo sapiens hypothetical gene supported by L39957; L43086; M85256; M95603; S56182; S73911; U07989, U79763; U62590; U62594; U6937; X58082; Z18127; AB006847; AB027436; AB027438; AF007572; AF035035; AF052535; AF099196; AF103384; AF103394;	
2330	AIVI_033463	Homo sapiens target of methylation-induced silencing 1 (TMS1) gene.	9E-14
2357	AF184072	complete cds	6.6

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Homo sapiens region containing TLS-associated serine-arginine	
		protein 1, TLS-associated serine-arginine protein 2, TLS-associated	
2250	7/3 f 0000 tm	serine-arginine protein 2; TLS-associated serine-arginine protein 2;	
2358	XM_032347	TLS-associated serine-arginine protein 1; TLS-associa>	0
2360	X54807	Human CYP2C8 gene for cytochrome P-450, 5' flank and exon 1	0.00003
2361	X67506	C.thermocellum ancA gene	0.0003
		Human DNA sequence from clone RP11-509D8 on chromosome 9,	0.034
2362	AL513350	complete sequence [Homo sapiens]	0.003
2363	AF020663	Homo sapiens myotubularin (MTM1) gene, promoter and exon 1	0.068
		Human chromosome 15q11-q13 putative DNA replication origin in	0,000
2364	U43626	the g-aminobutyric acid receptor b3 and a5 gene cluster	0.059
		Oryza sativa microsatellite MRG1415 containing (AT)X26, genomic	0,007
2365	AY019090	sequence	0.081
2366	U27586	Cytophaga heparina heparinase III (HepC) gene, complete cds	0.38
-		Homo sapiens similar to MYOSIN HEAVY CHAIN, CARDIAC	
		MUSCLE ALPHA ISOFORM (MYHC-ALPHA) (M. musculus)	
2367	XM_047192	(LOC92771), mRNA	e-133
		S.cerevisiae ATR1 gene conferring aminotriazole resistance, complete	
2368	M20319	cds	1.3
2369	XM 051036	Homo sapiens carbohydrate (keratan sulfate Gal-6) sulfotransferase 1	
2309	VM_021030	(CHST1), mRNA Macaca fascicularis brain cDNA clone:QmoA-12201, full insert	4.6
2370	AB062951	sequence	1.5
25.0	112002751	Pisum sativum clone MFRTPCR62 wound-inducible cytochrome P450	1,3
2371	U29335	mRNA, partial cds	0.41
		Ureaplasma urealyticum serovar 14 23S ribosomal RNA and 5S	0.41
		ribosomal RNA genes, complete sequence; and dihydrofolate	
2372	AF272630	reductase (fol A) gene, partial cds	0.061
		Mycoplasma fermentans orfD1 gene, Insertion sequence IS1630D,	
2373	AF179375	orfD2 gene	0.61
2374	X16876	Soybean ENOD2B gene for Ngm-75	0.96
2375	AF263831	Homo sapiens chromosome 11 DNaseI hypersensitive region	2.7
2376	AF157162	Syzygites megalocarpus 18S ribosomal RNA gene, partial sequence	5.4
2277	4 1200 500	Medicago truncatula partial mRNA for nodulin 25 (nod25 gene), clone 3	
2377	AJ308588 X02870	Bovine gene for cytokeratin VIb	4.6
2310	A02870		0.053
2379	AL590157	Human DNA sequence from clone RP13-420K18 on chromosome X, complete sequence [Homo sapiens]	417.09
2317	, ALJ 701J /	Plasmodium falciparum strain Dd2 chloroquine resistance marker	4E-08
2380	AF282242	protein gene, complete eds	4.2
2381	Z24745		
		A.thaliana 2S albumin gene isoforms 1 and 2, complete CDS's	0.6

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Human DNA sequence from clone RP1-14113 on chromosome	
2382	AL035418	22q13.1-13.33 Contains a GSS, complete sequence [Homo sapiens]	1.6
2383	Y17968	Gallus gallus mRNA for high mobility group 1 protein	1.5
2384	XM_034735	Homo sapiens similar to hypothetical protein FLJ21463 (H. sapiens) (LOC90883), mRNA	5.6
2385	AF001415	Arabidopsis thaliana 14-3-3-like protein GF14 upsilon (GRF5) gene, complete cds	1.3
2386		Homo sapiens retinol dehydrogenase 8 (all-trans) (RDH8), mRNA	1.9
2387	XM_003095	Homo sapiens zinc finger protein 288 (ZNF288), mRNA	1.8
2200	ATOMOS CO	Drosophila melanogaster genomic scaffold 142000013385449,	0.54
2388	AE003083	complete sequence	0.51
2200	ATOOMES	Drosophila melanogaster genomic scaffold 142000013385466,	0.014
2389	AE002798	complete sequence	0.018
2390	AL359506	Human DNA sequence from clone CTD-2007A7 on chromosome 20. Contains GSSs, complete sequence [Homo sapiens]	0.28
2370	7111333300	comania disas, complete sequence [fromo sapiens]	0.26
2391	U32726	Haemophilus influenzae Rd section 41 of 163 of the complete genome	0.3
2392	AB010468	Cyprinus carpio mRNA for CXC chemokine receptor-1, complete cds	0.9
2393	AE006886	Sulfolobus solfataricus section 245 of 272 of the complete genome	0.19
2394	M19810	Human apolipoprotein B-100 (apoB) gene, exon 4	0.43
2395	AB005548	Homo sapiens gene for squamous cell carcinoma antigen 2, partial cds	0.0004
2396	AJ318337	Plasmodium falciparum krueppell gene for Krueppel-like protein	0.62
2370	713510557	Human lung Clara cells 10 kda secretory protein (CC10) gene,	0.02
2397	U01102	satellite and Alu repeat sequences, complete eds	3.8
2377		Mus musculus 8 days embryo cDNA, RIKEN full-length enriched	3.0
2398	AK017766	library, clone:5730512J02, full insert sequence	1.7
2399	AE000032	Mycoplasma pneumoniae M129 section 48 of 63 of the complete genome	0.55
2400	AF333234	Homo sapiens receptor activator of nuclear factor kappa B ligand (RANKL) gene, promoter region	0.22
2401	BC010652	Homo sapiens, clone MGC:9753 IMAGE:3855206, mRNA, complete cds	1.8
2402	AK007856	Mus musculus 10 day old male pancreas cDNA, RIKEN full-length enriched library, clone:1810054D07, full insert sequence	2E-94
2403	AF172723	Homo sapiens integrin alpha 10 subunit (ITGA10) gene, exons 7 through 17 and partial cds	0.21
2404	NM 026570	Mus musculus glioma-amplified sequence-41 (Gas41-pending), mRNA	1.1
2405	U78770	Mus musculus spasmolytic polypeptide (mSP) gene, complete cds	1.8

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
2406	U26729	Human neprilysin gene, non-coding exon 3	0.92
		Lactococcus lactis subsp. lactis IL1403 section 108 of 218 of the	
2407	AE006346	complete genome	2
2400	. =======	Felis catus Niemann-Pick type C1 disease protein (NPC1) mRNA,	
2408	AF258783	complete cds	0.23
2400	4.000=200	Homo sapiens BAC clone RP11-332C7 from 7p11.2-p21, complete	
2409 2410	AC007280	sequence	0.39
2410	D13065	Mycoplasma capricolum mcs4 gene for MCS4 RNA	0.062
2411	4E100050	Drosophila pseudoobscura strain PSU606 period (per) gene, intron 4,	
2411	AF196058	partial sequence	1.5
2412	AT2370524	Arabidopsis thaliana Putative L-ascorbate peroxidase (AT4g09010)	[
2412	AF370534	mRNA, complete cds	1.8
2413	AF084197	Pongo pygmacus gamma-aminobutyric acid receptor A5 subunit	[
2413	Y17045	duplicated gene, 5'UTR region	0.00009
2414		Plasmodium falciparum gltS gene	0.063
2413	AL021177	Caenorhabditis elegans cosmid Y1A5A, complete sequence	0.22
2416	AJ238847	Drosophila melanogaster mRNA for drosophila dodeca-satellite protein 1 (DDP-1)	
2417	AB056333		0.22
2417	AF104821	Macaca fascicularis brain cDNA, clone:QflA-13486	e-126
2410	AF104621	Danio rerio Hlx2 gene, promoter region	0.056
ĺ		Neospora caninum ycf24 protein (ycf24) gene, partial cds; DNA	
		dependent RNA polymerase beta subunit (rpoB) gene, complete cds;	
2419	AF138960	and DNA dependent RNA polymerase beta subunit' (rpoCl) gene, partial cds, plastid genes for plastid products	
2420	AF338712	Emeus crassus mitochondrion, partial genome	0.21
2421	Z82625	R. prowazekii genomic DNA fragment (clone A396F)	1.3
2121	202025	In prowazekii genomie DNA nagineni (cione A396F)	0.46
2422	XM 015945	Homo sapiens hypothetical protein FLJ13842 (FLJ13842), mRNA	6E-62
2423	X57010	Human COL2A1 gene for collagen II alpha 1 chain, exons E2-E15	0.63
2424	AF283525	Homo sapiens FIP2 gene, exons 11, 12 and 13	4E-71
		Mus musculus adult male hippocampus cDNA, RIKEN full-length	
2425	AK013530	enriched library, clone:2900012M01, full insert sequence	5.3
2426	D86077	Homo sapiens DNA for cyclin G, partial cds	0.02
2427	X94208	H.sapiens TPR gene (1121bp)	0.19
		Nicotiana tabacum Avr9/Cf-9 rapidly elicited protein 4 (ACRE4)	
2428	AF211528	mRNA, complete cds	0,22
1			
2430	L35688	Homo sapiens (subclone H9 10_d10 from P1 LBL#3) DNA sequence	5E-08
ĺ		Cavia porcellus seminal vesicle secretory protein SVP-1/-3/-4 gene,	
2431	U59711	complete cds	0.14
2432	AK023305	Homo sapiens cDNA FLJ13243 fis, clone OVARC1000622	0.002
2433	AB048868	Macaca fascicularis brain cDNA, clone:QnpA-10055	0.044
		Homo sapiens clone C13 immunoglobulin heavy chain variable region	
2434	AF167585	gene, partial cds	0.13
2435	AB013796	Ceratopteris richardii mRNA for CRHB6, complete cds	0.024

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
2436	AF071481	Homo sapiens potassium channel (KCNQ3) gene, exon 4	1.9
		Plasmodium falciparum strain Dd2 chloroquine resistance marker	
2437	AF282242	protein gene, complete cds	0.22
		Oryza sativa microsatellite MRG0632 containing (AT)X14, genomic	
2438	AY018307	sequence	0.073
2439	Z74273	S.cerevisiae chromosome IV reading frame ORF YDL225w	0.028
2440	A3700105#	Oryza sativa microsatellite MRG3582 containing (TA)X22, genomic	
2440 2441	AY021257	sequence	0.018
2441	XM_012164	Homo sapiens STE20-like kinase (JIK), mRNA	0.27
	TD - 00	Homo sapiens hypothetical protein DKFZp564B1023	
2442	XM_036587	(DKFZP564B1023), mRNA	0.008
2442	47 110540	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen	
2443	AL112743	deprivation	0.069
2444	AB020575	Conocephalum conicum psbA gene, complete cds, strain J type	0.074
2445	DC001550	Homo sapiens, Similar to glutaminyl-tRNA synthetase, clone	
2443	BC001772	MGC:1525 IMAGE:3543728, mRNA, complete cds	5.5
2446	4E000110	Drosophila montana clone mon 10 transposon Penelope gene	
2440	AF289113	sequence	0.064
2447	VA 01 5006	TY	
2447	XM_015806 AY047559	Homo sapiens hypothetical protein FLJ22621 (FLJ22621), mRNA Drosophila melanogaster GH07383 full length cDNA	0.92
2448	XM 052662	Homo sapiens amino-terminal enhancer of split (AES), mRNA	
2449	AW_032062	Human DNA sequence from clone RP11-446N1 on chromosome 6,	0.59
2450	AL138723	complete sequence [Homo sapiens]	0.000
2430	AL/130723	Dermatobia hominis strain Ponta Grossa tRNA-Ile gene, partial	0.008
		sequence; D-loop, complete sequence; and 12S ribosomal RNA,	
2451	AF151389	partial sequence; mitochondrial genes for mitochondrial products	0.46
2431	AF131369	partial sequence, intochondrial genes for intochondrial products	0.40
2452	NM 018123	Homo sapiens hypothetical protein FLJ10517 (FLJ10517), mRNA	0.21
2432	14M_010123	Homo sapiens hypothetical gene supported by AL359651	0.21
2453	XM_042803	(LOC92091), mRNA	1.9
2100	7111_012005	Plasmodium falciparum chromosome 2, section 34 of 73 of the	1.9
2454	AE001397	complete sequence	5
2455	AF017996	Mus musculus lysosomal acid lipase gene, promoter region	0.022
2456	XM 028409	Homo sapiens KIAA1871 protein (KIAA1871), mRNA	0.0003
2457	XM 051692	Homo sapiens KIAA1705 protein (KIAA1705), mRNA	0,0003
2458	U00037	Caenorhabditis elegans cosmid T20H4	0.073
2459	AL138748	Human DNA sequence from clone RP4-732E19 on chromosome Xq21.1-21.33 Contains GSSs, complete sequence [Homo sapiens] Homo sapiens transcriptional repressor p54 gene, promoter region and	0.2
2460	AF052744	partial cds	1.8
2461	XM 007006		1.8
2462	AF303046	Homo sapiens prostinogen mRNA, complete cds	0.44
2463	AK023562	Homo sapiens cDNA FLJ13500 ffs, clone PLACE1004693	e-116
2403	11023302	LIGHTO SAPIGHS GOVER PLACE 1004093	6-110

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
		Aspergillus fumigatus aureobasidin-resistance protein (aur1) mRNA,	1 VALUE
2464	AF076692	complete cds	3.2
		Glossina morsitans morsitans thrombin inhibitor precursor (TTI)	
2465	AF054616	mRNA, complete cds	0.44
2466	754147	Human DNA sequence from cosmid L129H7, Huntington's Disease	
2400	Z54147	Region, chromosome 4p16.3 contains CpG island Homo sapiens glutamate receptor, ionotropic, AMPA 1 (GRIA1),	4E-09
2467	XM 040290	mRNA (GRIA1),	4
2107	11.12_070250	Rattus norvegicus dynein-associated protein RKM23 (km23) mRNA.	
2468	AY026512	complete cds	1.5
2469	AY033091	Plasmodium berghei Pbs36 gene, complete cds	0.0008
2470	XM_051572	Homo sapiens heat shock transcription factor 2 (HSF2), mRNA	2E-42
2471	AF287003	Floriceps minacanthus 18S ribosomal RNA gene, complete sequence	1.8
2472	X99872	Sulfolobus acidocaldarius pyrB gene	0.2
1		Bombina orientalis SAP bombesin preprohormone mRNA, complete	
2473	U49451	cds	0.56
2474	AC024819	Caenorhabditis elegans cosmid Y55B1AL, complete sequence	6.9
		Litoria subglandulosa strain SAMA_R51051 NADH dehydrogenase	
	4 Taga (00	subunit 4 (ND4) gene, partial cds; mitochondrial gene for	
2475	AF282602	mitochondrial product	0.073
2476	AL591343	Human DNA sequence from clone RP11-260H5 on chromosome 6,	
24/0	AL391343	complete sequence [Homo sapiens] Human DNA sequence from cosmid L174G8, Huntington's Disease	0.24
2477	Z69375	Region, chromosome 4p16.3	2F 16
	205575	rogion, enrollosome 4p10.5	3E-16
2478	U32174	Dictyostelium discoideum spore lysis A (splA) gene, complete cds	0.25
2479	XM 045095	Homo sapiens KIAA1320 protein (KIAA1320), mRNA	0.53
2480	AY035224	Gallus gallus cardiac muscle factor 1 gene, partial cds	0.78
		Penicillium urticae mitochondrial I-rRNA (large rRNA) gene and its	
2481	D14567	flanking region	0.25
		HIV-1 strain 96CG38.10 from Republic of the Congo, envelope	
2482	AF119218	glycoprotein, C2-C4 region (env) gene, partial cds	0.002
2483	AF386963	Arabidopsis thaliana Unknown protein mRNA, complete cds	0.057
2484	Z19595	C.elegans her-1 gene	0.19
		Lactococcus lactis subsp. lactis IL1403 section 103 of 218 of the	
2485	AE006341	complete genome	6.1
		Human DNA sequence from clone RP11-469L10 on chromosome 6,	
2486	AL583831	complete sequence [Homo sapiens]	0.23
2487	NC_001617	Human rhinovirus 89	0.078
2488	X65859	H.sapiens pseudogene for the low affinity IL-8 receptor	5.7
2489	AE007508	Strentococome pnormanics restion 101 of 104 of the complete	0.62
2490	AF199367	Streptococcus pneumoniae section 191 of 194 of the complete genome Entamoeba histolytica cdc48-like protein gene, complete cds	0.63
2470	131.13230/	Pseudonaja textilis tissue-type liver phospholipase A2 inhibitor beta	0,30
2491	AF211160	subunit isoform PTI-2B mRNA, complete cds	2.2
	211100	sucum notom r 11-2D micror, complete cus	2.2

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	
140	ACCESSIV	Methanobacterium thermoautotrophicum from bases 206991 to	P VALUE
2492	AE000813		
2492	AE000613	217929 (section 19 of 148) of the complete genome	5.5
2493	U67604	Methanococcus jannaschii section 146 of 150 of the complete genome	1
		Pichia canadensis mitochondrial gene for NADH dehydrogenase	
2494	D16253	subunit 5, complete cds	0.006
2495	NC_001530	Human papillomavirus type 47, complete genome	5.4
ì i		Pseudavenzoaria ochropodis from host Tringa ochropus 16S	
		ribosomal RNA gene, partial sequence; mitochondrial gene for	
2496	AF286437	mitochondrial product	0.4
2497	XM_006069	Homo sapiens wee1+ (S. pombe) homolog (WEE1), mRNA	3E-26
		Rana boylii NADH dehydrogenase subunit I (ND1) gene, partial cds:	
		tRNA-lle, tRNA-Gln, and tRNA-Met genes, complete sequence;	
		NADH dehydrogenase subunit II (ND2) gene, complete cds; tRNA-	
		Trp, tRNA-Ala, tRNA-Asn, tRNA-Cys, and tRNA-Tyr genes,	
2498	AF314019	complete se>	1.4
2499	AJ314911	Dictyostelium discoideum DrnA gene for putative RNaseIII, exons 1-3	0.51
		Vesicular stomatitis virus (strain 86-GM-B) glycoprotein gene.	
2500	M35228	complete cds	0.45
2501	AB021290	Homo sapiens gene for chondromodulin-1, promoter and partial cds	0.72
2502	AJ307807	Astrammina rara partial 18S rRNA gene, isolate A44	3.5
2503	AB005148	Bos taurus mRNA for IL-1 receptor antagonist, complete cds	0.84
		Drosophila melanogaster genomic scaffold 142000013385743,	
2504	AE002687	complete sequence	0.23
		Caenorhabditis elegans beta chain spectrin homolog Sma1 (sma1)	
2505	AF053496	mRNA, complete cds	5.9
,		E.coli IncFV plasmid pED208 oriT region with genes traM, traJ and	
2506	X59611	traY (part.)	0.043
		Homo sapiens phospholipase A2, group IVA (cytosolic, calcium-	
2507	XM_051897	dependent) (PLA2G4A), mRNA	3E-66
2508	AJ132890	Bos taurus mRNA for acetyl-CoA-carboxylase	0.18
2509	Y13537	Homo sapiens LAG-3 gene, 3' region	1.7
2510	AE002149	Ureaplasma urealyticum section 50 of 59 of the complete genome	0.027
		Influenza C virus C/Johannesburg/1/66 segment 5 nucleoprotein (NP)	
2511	AF170573	gene, partial cds	2.4
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
2512	AK016507	library, clone:4931439C15, full insert sequence	6.8
		· ·	
1		Human DNA sequence from clone RP11-209D8 on chromosome 6.	
		Contains ESTs and GSSs. Contains part of the COL12A1 gene for	
2513	AL354664	collagen type XII, alpha 1, complete sequence [Homo sapiens]	0.082
		Pisum sativum PsEXT mRNA for xyloglucan endotransglycosylase,	
2514	AB042531	complete cds	2E-12
			

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
2515	AL590371	Human DNA sequence from clone RP11-295H23 on chromosome 6, complete sequence [Homo sapiens]	1.5
2516	AF334780	Homo sapiens pregnancy-induced growth inhibitor OKL38 gene, partial cds	0.8
2517	AF362371	Dictyostelium discoideum histidine kinase DhkI (dhkI) gene, complete cds	0.021
2518	NM_012608	Rattus norvegicus Membrane metallo-endopeptidase (neutral endopeptidase/enkephalinase) (Mme), mRNA	2
2519	BC007173	Mus musculus, clone IMAGE:3493127, mRNA, partial cds	5E-53
2520	Y13870	Homo sapiens mRNA containing (CAG)6 repeat, clone CZ-CAG-12	0.06
		Homo sapiens hyaluronidase-like pseudogene 1 (HYALP1), partial	
2521	AF051769	sequence	0.009
		Dictyostelium discoideum Rac1A (rac1A) gene, complete cds; and	
2522	AF309947	unknown gene	0.77
2523	AF128392	Arabidopsis thaliana BAC F15P23	0,22
		Mus musculus adult male spinal cord cDNA, RIKEN full-length	
2524	AK020739	enriched library, clone: A330106L17, full insert sequence	7.2
2525	Y09472	B.taurus gene encoding preprododecapeptide	0.24
		Rat DNA for 3'UTR of skeletal muscle sodium channel, partial	
2526	D87922	sequence	0.4
2527	U70865	Carassius auratus CCK8 precursor, mRNA, complete cds	0.67
		Homo sapiens FLI1 gene for ERGB transcription fuctor, intron 4 and	
2528	AB012624	partial cds	0.054
2529	AF180492	Crocuta crocuta microsatellite Ccr12 sequence	0.083
		Human DNA sequence from clone 536P6 on chromosome 22.	
2530	AL078640	Contains an EST, complete sequence [Homo sapiens]	2E-11
2531	Z63184	H.sapiens CpG island DNA genomic Msel fragment, clone 7al, forward read cpg7al.ftld	6E-22
		Homo sapiens mRNA; cDNA DKFZp761M2223 (from clone	
2532	AL162055	DKFZp761M2223)	0.022
2533	AE001392	Plasmodium falciparum chromosome 2, section 29 of 73 of the complete sequence	6.5
		Sporopachydermia cereana var. D77-321B internal transcribed spacer	
		1, 5.8S ribosomal RNA gene and internal transcribed spacer 2,	
		complete sequence; and large subunit ribosomal RNA gene, partial	
2535	AF202903	sequence	0.14
		Borrelia burgdorferi P35 antigen protein gene, and 7.5 kDa	
2536	U59487	lipoprotein gene, complete cds	0.029
2537	XM_030999	Homo sapiens KIAA0444 protein (KIAA0444), mRNA	0.012
2538	AF302838	Mus musculus putative tumor suppressor LEU2 gene, exons 1 and 2	0,63
		Bresiliidae gen. sp. LSP21-1 cytochrome oxidase I gene, partial cds;	
2539	AF125423	mitochendrial gene for mitochendrial product	0.66
		Homo sapiens genomic DNA, chromosome 21q21.1-q21.2	
2540	AP001348	clone:f39G8, LL56-APP region, complete sequence	0.42

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	-	Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEO ID			
NO	ACCESSN	DESCRIP	P VALUE
2541	XM_028324	Homo sapiens hypothetical protein FLJ14195; KIAA1714 protein (FLJ14195), mRNA	8E-98
2542	AF238860	Streptococcus mutans mutacin I biosynthesis gene cluster, partial sequence	4.6
2543	AE005391	Escherichia coli O157:H7 EDL933 genome, contig 3 of 3, section 10 of 290	1.6
2544	AK024518	Homo sapiens cDNA: FLJ20865 fis, clone ADKA01850	0.06
		Homo sapiens cDNA: FLJ22942 fis, clone KAT08170, highly similar	
2545	AK026595	to HUMMHBW62 Homo sapiens (clone pMF18) MHC class I HLA- Bw62 mRNA	4E-39
2546	AL449163	Human DNA sequence from clone RP11-533E16 on chromosome 6, complete sequence [Homo sapiens]	2.3
2547	AJ305053	Caprine arthritis-encephalitis virus proviral partial pol gene for Pol protein (reverse transcriptase), isolate CAEV Br/UFRGS-2/C767	0,69
2548	AF283067	Homo sapiens clone 3ptel_c11bt3 sequence	1E-41
2549	AF286431	Bychovskiata intermedia from host Charadrius leschenaulti 16S ribosomal RNA gene, partial sequence; mitochondrial gene for mitochondrial product	0.94
2550	L35232	Schizosaccharomyces pombe deoxyribonuclease (exo2) gene, exons 1-3, complete cds	0.93
2551	AF319044	Methanosarcina acetivorans strain C2A MtaF (mtaF) and MtaG (mtaG) genes, complete cds	0.034
2552	XM_018334	Homo sapiens hypothetical protein FLI22418 (FLJ22418), mRNA	0.013
2553	U35862	Visna virus clone 678 env3.2 envelope polyprotein (env) gene, partial cds	0.17
2554	M20005	N.crassa conidation-specific protein genes con-10 (complete cds.) and con-13 (3' flank)	0.079
2555	AB050531	Macaca fascicularis brain cDNA, clone:QnpA-20907	1.4
2556	AB009693	Mus musculus gene for mafG, complete cds	5.6
2557	U67593	Methanococcus jannaschii section 135 of 150 of the complete genome	0.024
2558	XM_028292	Homo sapiens x 009 protein (MDS009), mRNA	4
0550	VICARAO		
2559	U67578 AE001759	Methanococcus jannaschii section 120 of 150 of the complete genome Thermotoga maritima section 71 of 136 of the complete genome	1.8
2300	AE001739	Phascolion strombi homeodomain transcription factor Gsx (Gsx)	1.9
2561	AF363230	gene, partial cds	0,59
2562	NM_019211	Rattus norvegicus RAS guanyl releasing protein 1 (Rasgrp), mRNA	2.5
2563	U60070	Solanum tuberosum disease resistance homolog (St121) gene, partial eds	0,68
2564	AE006593	Streptococcus pyogenes M1 GAS strain SF370, section 122 of 167 of the complete genome	0,062

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Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE Wilkesia gymnoxiphium ASAPETALA1-A (ASAP1-A) gene, exons 3 2565 AF147231 through 8, partial cds 0.026 Mus musculus adult male cecum cDNA, RIKEN full-length enriched 2566 AK018656 library, clone:9130402C12, full insert sequence 0.74 2567 X65859 H. sapiens pseudogene for the low affinity IL-8 receptor 5.4 Homo sapiens similar to KIAA1361 protein (H. sapiens) (LOC90315), 2568 XM 030855 mRNA 0.65 2569 XM 031726 Homo sapiens hypothetical protein FLJ12577 (FLJ12577), mRNA 2.1 Caenorhabditis elegans cosmid F56H11, complete sequence 2570 Z68749 0.025 Mycoplasma pneumoniae M129 section 60 of 63 of the complete 2571 AE000020 genome 0.17 Oryctolagus cuniculus beta-globin-like gene cluster locus control 2572 U63091 region 0.13 2573 BC001339 Homo sapiens, clone IMAGE:3461987, mRNA, partial cds 1E-31 2574 U86962 Dictvostelium discoideum MigA (migA) gene, complete cds 0.9 Oryza sativa microsatellite MRG3954 containing (TA)X30, genomic 2575 AY021629 sequence 6.5 Macaca fascicularis brain cDNA clone:OtrA-13349, full insert 2576 AB063077 semience 0.59 2577 XM 044332 Homo sapiens piwi (Drosophila)-like 1 (PIWIL1), mRNA 4E-28 Z97185 S.pombe chromosome I cosmid c16A10 2578 1.5 2579 L43603 Gallus gallus Na+/K+-ATPase alpha-1 subunit gene, promoter region 0.7 2580 AC016645 Homo sapiens chromosome 5 clone RP11-54A24, complete sequence 0.0001 Plasmodium falciparum chromosome 2, section 45 of 73 of the 2581 AE001408 complete sequence 0.25 Ischmura prognata cytochrome b (cytb) gene, mitochondrial gene 2582 AF067699 encoding mitochondrial protein, partial cds 0.45 2583 AC026061 Homo sapiens BAC clone RP11-223K9 from Y, complete sequence 2E-15 2584 Y13537 Homo sapiens LAG-3 gene, 3' region 0.24 Caulobacter crescentus section 303 of 359 of the complete genome 2585 AE005977 5.5 Angiostrongylus cantonensis clone G-2 retrotransposable element 2586 U13190 0.077 2587 AF364550 Drosophila melanogaster transposon cruiser, complete sequence 0.07 Homo sapiens CD36 antigen (collagen type I receptor, 2588 XM 034146 thrombospondin receptor) (CD36), mRNA 0.62 Gonioctena viminalis 12S ribosomal RNA gene, mitochondrial gene 2589 AF014580 for mitochondrial RNA, partial sequence 2.1 Homo sapiens hypothetical protein DKFZp434G131 XM_042643 2590 (DKFZP434G131), mRNA 0.68 Homo sapiens, ribosomal protein \$16, clone MGC:10931 2591 BC004324 IMAGE:3628799, mRNA, complete cds 0.006

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO 2592	ACCESSN U42430	DESCRIP	P VALUE
2592	U42430	Mesocricetus auratus CD36 mRNA, complete cds	0.71
		Saccharomyces cerevisiae Mvp1p (MVP1) gene, complete cds	4.7
2594	AB047954	Macaca fascicularis brain cDNA, clone:QnpA-12630	0.65
	AF261697	Homo sapiens clone 17ptel_c39t3 sequence	0.024
2596	AF162598	Drosophila virilis strain w159 zeste protein (z) gene, partial cds	0.063
		Peromyscus sejugis ND3 and ND4L genes, complete cds, tRNA(arg)	
		gene, complete sequence, tRNA(gly) gene, partial sequence, and ND4	
		gene, partial cds, mitochondrial genes encoding mitochondrial	
2597	U40255	products	0.67
		H.sapiens CpG island DNA genomic Mse1 fragment, clone 7h10,	
2598	Z63251	forward read cpg7h10.fla	0.003
	******	Chelonus sp. 16S mitochondrial ribosomal RNA, mitochondrial gene,	
2599	U68150	partial sequence	0.003
2600	AF138281	Arabidopsis thaliana phospholipase D-gamma-2 mRNA, complete cds	0.071
		Mus musculus nitrilase homolog 1 (Nit1) gene, alternatively spliced	
2601	AF069985	product, complete cds	0.39
2602	M83134	Human germline IgH chain (hv3005) V3-region	0.35
		Trama rara 12S small subunit ribosomal RNA gene, partial sequence;	
		tRNA-Val gene, complete sequence; and 16S large subunit ribosomal	
		RNA gene, partial sequence; mitochondrial genes for mitochondrial	
2603	AF275245	products	0.59
2604	X93302	S.cerevisiae mRNA for Msn5 protein	0.05
l 1		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
2605	AK015536	library, clone:4930470H18, full insert sequence	0.7
2606	AC087735	Caenorhabditis elegans cosmid Y35H6, complete sequence	0.022
2607	AJ271002	Mus musculus TFF1/pS2 gene for Trefoil Factor1/pS2, exons 1-3	0.73
		Mus musculus potassium inwardly-rectifying channel, subfamily J,	
2608	NM_019664	member 15 (Kcnj15), mRNA	5.8
		Human DNA sequence from clone RP11-162H13 on chromosome 6	
2609	AL450330	Contains GSSs, complete sequence [Homo sapiens]	0.25
1		H. sapiens (D1S514) DNA segment containing (CA) repeat; clone	
2610	Z24671	AFMa151za5; single read	0.007
2611	AE000773	Aquifex aeolicus section 105 of 109 of the complete genome	0.34
		E.histolytica mRNA for 170kDa subunit of galactose/N-	
2612	X93508	acetylgalactosamine binding lectin	0.054
		Mus musculus 18 days embryo cDNA, RIKEN full-length enriched	
2613	AK003780	library, clone:1110018I21, full insert sequence	0.15
		Homo sapiens (subclone 1_c12 from P1 H54) DNA sequence,	
2614	L81687	complete sequence	5.2
2615	M10090	Human myoglobin gene, exon 1	0.13
2616	NC_002670	Bacteriophage bIL311, complete genome	0.061
2617	AB016259	Oryzias latipes gene for cardiac muscle actin, complete cds	0.19
2618	U67578	Methanococcus jannaschii section 120 of 150 of the complete genome	2

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID	1 aarami		
NO	ACCESSN	DESCRIP	P VALUE
2619	AF186258	Homo sapiens sulfotransferase 1C1 (SULT1C1) gene, exons 2 and 3	0.002
2620	X79777	P.furiosus aor, cmo and ado-hcy genes	6.1
2621	XM_042013	Homo sapiens cell division cycle 34 (CDC34), mRNA	1.5
2622	AB044879	Equus caballus DNA, microsatellite TKY379	0.18
2623	AF117951	Mus musculus lysyl oxidase-like protein 2 mRNA, partial cds	0.015
		Dictyostelium discoideum RacB (racB) gene, partial cds; and	
2624	AF310887	unknown gene	0.17
2625	XM_034353	Homo sapiens KIAA1500 protein (KIAA1500), mRNA	4E-89
2626	AK021859	Homo sapiens cDNA FLJ11797 fis, clone HEMBA1006182	0
2627	AF280894	Pongo pygmaeus clone 3458 chromosome Xq genomic sequence	0.0002
		Plasmodium falciparum chromosome 2, section 33 of 73 of the	
2628	AE001396	complete sequence	0.074
		Homo sapiens similar to RIKEN cDNA 2610307I21 gene;	
		hypothetical protein, MNCb-4273 (M. musculus) (LOC92606),	l
2629	XM_046097	mRNA	0.58
2630	X02438	Leishmania tarentolae maxicircle DNA fragment	· 6.2
		Bombus terrestris 16S large subunit ribosomal RNA gene, partial	
2631	AF181582	sequence	0.2
2632	M34149	S. cerevisiae mitochondrial gap 12 sequence	0.64
		Trioxys angelicae partial mitochondrial ATPase 6 gene for ATPase 6	
2633	AJ400612	subunit	0.65
2634	AB021290	Homo sapiens gene for chondromodulin-1, promoter and partial cds	0.75
2635	AJ133756	Drosophila melanogaster mRNA for jim zinc finger protein	0.028
		Human DNA sequence from clone RP5-1140M3 on chromosome 20.	İ
2636	AL121589	Contains GSSs, complete sequence [Homo sapiens]	0.002
2637	XM_040430	Homo sapiens hypothetical protein FLJ10599 (FLJ10599), mRNA	0.24
2638	AB015484	Dugesia japonica mRNA for myosin heavy chain, complete eds	0.2
2639	XM_008161	Homo sapiens hypothetical protein FLJ20345 (FLJ20345), mRNA	0.075
		Mus musculus gamma-aminobutyric acid receptor rho2 subunit	
2640	AF024621	mRNA, complete cds	6.8
2641	L78777	Homo sapiens (subclone 2_b8 from P1 H49) DNA sequence	8E-11
2642	AE001028	Archaeoglobus fulgidus section 79 of 172 of the complete genome	2
2643	AB013796	Ceratopteris richardii mRNA for CRHB6, complete cds	0.028
		Chlorella ellipsoidea chloroplast 16S-23S DNA spacer and ORF1 to	
2644	X03848	ORF4 DNA	0.086
		B. aphidicola trpE and trpG genes for anthranilate synthase,	
2645	Z21938	components I and II	0.74
		,	
2646	XM_039805	Homo sapiens hypothetical protein MGC3040 (MGC3040), mRNA	2.1

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Stenomesson variegatum ribulose 1,5-bisphosphate carboxylase large	
		subunit (rbcL) gene, partial cds, chloroplast gene for chloroplast	
2704	AF116983	product	6.4
2706	XM_051524	Homo sapiens archain 1 (ARCN1), mRNA	0.66
		Homo sapiens lipocalin-1 interacting membrane receptor (LIMR)	
2707	AF351620	gene, complete cds	0.084
		Homo sapiens aryl-hydrocarbon interacting protein-like 1 (AIPL1)	
2708	AF180472	gene, complete cds	5.9
		Ictalurus punctatus clone Icpu-UA/3 MHC class I antigen gene,	
2709	AY008848	complete cds	0.025
		Mus musculus vesicle transport through interaction with t-SNAREs 1	
2710	NM_016862	homolog (Vti1-pending), mRNA	0.65
2711	AE004023	Xylella fastidiosa 9a5c, section 169 of 229 of the complete genome	3.9
2712	XM_037239	Homo sapiens actin, beta (ACTB), mRNA	1E-49
		Plasmodium falciparum chromosome 2, section 41 of 73 of the	
2713	AE001404	complete sequence	0.7
2714	XM_050114	Homo sapiens TERA protein (TERA), mRNA	5.9
2715	AK001082	Homo sapiens cDNA FLJ10220 fis, clone HEMBA1007342	0.56
2716	AF380155	Ctenopharyngodon idella c-Fos (c-fos) mRNA, complete cds	3.2
		Cordyceps sp. 97003 mitochondrial DNA for small subunit ribosomal	
2717	AB027352	RNA	0.47
		Lactococcus lactis gadB gene, untranslated by upstream frameshift,	
2718	AB033220	strain:01-1	1.9
		Mus musculus B6/CBAF1J peroxisome proliferator activated receptor	
2719	U01841	gamma (ppar) mRNA, complete cds	0.009
2720	AL023814	Caenorhabditis elegans cosmid H08J19, complete sequence	0.021
2721	XM_047737	Homo sapiens ubiquitin specific protease 25 (USP25), mRNA	5E-56
		Takifugu rubripes pericentriolar material 1 protein (PCM1) mRNA,	
2723	AF389403	complete cds	1.9
		Lactococcus lactis subsp. lactis IL1403 section 131 of 218 of the	
2724	AE006369	complete genome	0.21
ll		Mesembryanthemum crystallinum potassium transporter HAK2p	
2726	AF367865	(HAK2) mRNA, complete cds	0.39
2727	XM_043040	Homo sapiens hypothetical protein MGC2718 (MGC2718), mRNA	1E-09
		Plasmodium falciparum chromosome 2, section 29 of 73 of the	
2728	AE001392	complete sequence	0.0000004
2729	U31371	Bovine herpesvirus 4 1.7-kb late RNA (L1.7) mRNA, repeat region	1.9
l l		Libellula flavida 16S ribosomal RNA gene, partial sequence;	
2730	AF195728	mitochondrial gene for mitochondrial product	0.025
2731	AK024658	Homo sapiens cDNA: FLJ21005 fis, clone CAE03745	0.64
2732	AK025613	Homo sapiens cDNA: FLJ21960 fis, clone HEP05517	1.4
2733	L11016	Homo sapiens lymphotoxin-beta gene, complete cds	0,0001
2734	Y14422	M.musculus DNA for retinal protein	1.2

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
2735	X83757	E.histolytica 170 kDa lectin gene, 5' region	0.64
		Mus musculus 8 days embryo cDNA, RIKEN full-length enriched	
2736	AK017535	library, clone:5730409K12, full insert sequence	0.64
2737	AY047509	Drosophila melanogaster GH01721 full length cDNA	0.21 .
	17010100	Homo sapiens gene for T-cell nuclear receptor NOT (Nurr1), complete	
2738 2739	AB019433 U58731	cds	0.024
2/39	U38/31	Caenorhabditis elegans cosmid W09B12	0.61
1			
2740	U35620	Nicotiana tabacum ribulose-1,5 bisphosphate carboxylase/oxygenase	
2/40	U33620	large subunit N-methyltransferase (rbcmtT) mRNA, complete cds	1
2741	AF181720	Homo sapiens RU2AS (RU2) gene, complete cds; and RU2S (RU2) gene, partial cds	
2/41	AF161720	Streptococcus pyogenes M1 GAS strain SF370, section 89 of 167 of	5.4
2742	AE006560	the complete genome	0.04
2/42	AE000300	Gonioctena pallida haplotype 61 mitochondrial control region, partial	0.24
2743	AF346287	sequence	5
2744	L05617	Dictyostelium purpureum (Dpp5) DNA sequence, repeat region	0.021
2/44	103017	Mus musculus adult male corpus striatum cDNA, RIKEN full-length	0.021
2745	AK021104	enriched library, clone:C030026M15, full insert sequence	1
2743	7111021104	Romanomermis culicivorax mitochondrial NADH dehydrogenase	1
		subunits 6 and 3 (ND6, ND3) cytochrome P450-like protein genes,	
2746	L08174	complete CDS	1.6
2747	AB006130	Mesocricetus auratus mRNA for haptoglobin, complete cds	0.24
2748	AK001448	Homo sapiens cDNA FLJ10586 fis, clone NT2RP2003986	0,0008
		The state of the s	0.0000
2749	M95192	Drosophila melanogaster hairless protein (H) mRNA, complete cds	3.6
		Nicotiana sylvestris NySS41 and partial NySS42 genes for RUBISCO	
2750	X53426	small subunit (SSU)	0.63
		Methanobacterium thermoautotrophicum plasmid pFV1, complete	
2751	NC 001336	sequence	0.074
2752	NM 011595	Mus musculus tissue inhibitor of metalloproteinase 3 (Timp3), mRNA	0.012
2753	AF299340	Homo sapiens CD164 gene, complete cds, alternatively spliced	0.074
2754	AL121765	S.pombe chromosome I cosmid c1786	0.025
		Trichaptum abietimum mitochondrial small subunit ribosomal RNA,	
2755	U27078	mitochondrial gene, partial sequence	0.44
		Mycoplasma hyopneumoniae strain ISU 232 YX1 (yx1) gene, partial	
l l		cds; p110 membrane protein precursor, gene, complete cds; and YX2	
2756	AF279292	(yx2) gene, partial cds	0.024
		Human DNA sequence from clone RP11-295H23 on chromosome 6,	
2757	AL590371	complete sequence [Homo sapiens]	0.61
2758	XM_041904	Homo sapiens cathepsin S (CTSS), mRNA	6
1		Gallus domesticus DNA for the terminal heterochromatic region of	
2759	D63170	the Z chromosome	1.4
		Mus musculus RIKEN cDNA 2310008H09 gene (2310008H09Rik),	
2760	NM_023197	mRNA	0.025

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
2761	BC007150	Mus musculus, dystroglycan 1, clone MGC:6651 IMAGE:3496914, mRNA, complete cds	0.18
2762	BC000480	Homo sapiens, amyloid beta precursor protein-binding protein 1, 59kD, clone MGC:8690 IMAGE:2964638, mRNA, complete cds	8E-65
2764	XM_036683	Homo sapiens a disintegrin-like and metalloprotease (reprolysin type) with thrombospondin type 1 motif, 3 (ADAMTS3), mRNA	3E-46
2765	AJ005581	Encephalitozoon cuniculi complete rDNA unit (16S, 5.8S, 23S, ITS1 and IGS)	0.42
2766	AE005227	Escherichia coli O157:H7 EDL933 genome, contig 1 of 3, section 51 of 155	0.007
2767	XM_048172	Homo sapiens hypothetical protein DKFZp434F1017 (DKFZP434F1017), mRNA	3.8
2769	AF129087	Medicago sativa mitogen-activated protein kinase homologue (TDY1) gene, complete cds	0.66
2770	XM_008176		5.2
2771	AC024865	Caenorhabditis elegans cosmid Y74C10AL, complete sequence	0.039
2772	XM_036164	Homo sapiens RAN binding protein 2 (RANBP2), mRNA	1E-59
2773	AL023816	Caenorhabditis elegans cosmid T05G11, complete sequence	0.64
2774	XM_004980	Homo sapiens cystic fibrosis transmembrane conductance regulator, ATP-binding cassette (sub-family C, member 7) (CFTR), mRNA Homo sapiens hypothetical gene supported by AK023337	0.016
2775	XM_032209	(LOC90509), mRNA	0.018
2776	Z69650	Human DNA sequence from cosmid L69F7B, Huntington's Disease Region, chromosome 4p16.3 contains Huntington Disease (HD) gene	0.64
2777	AJ305053	Caprine arthritis-encephalitis virus proviral partial pol gene for Pol protein (reverse transcriptase), isolate CAEV Br/UFRGS-2/C767	0.46
2778	AK017531	Mus musculus 8 days embryo cDNA, RIKEN full-length enriched library, clone:5730409F18, full insert sequence	0.67
2779	AF028784	Rattus norvegicus glial fibrillary acidic proteins alpha and delta (GFAP) gene, alternatively spliced products, complete eds	0.11
2780	AF101236	Naja sputatrix neutral phospholipase A2 (NPLA2) gene, complete cds	0.054
2781	U71280	Mesocricetus auratus cytochrome P450 aldosterone synthase (CYP11B2) gene, complete cds	0.023
2782	X80007	P.sativum gene for chalcone synthase	0.15
2783	AF171040	Anopheles funestus clone AF10 microsatellite sequence	2
2784	XM_011464	Homo sapiens hypothetical protein FLJ20170 (FLJ20170), mRNA	0.21
2785	Y18929	Homo sapiens wild type AF-4/FEL gene, part of intron 3, individual PB	3
2786	L81581	Homo sapiens (subclone 4_a4 from P1 H11) DNA sequence, complete sequence	0.024

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Rattus norvegicus ATP-stimulated glucocorticoid-receptor	
2787	NM_024381	translocaton promoter (Gyk), mRNA	0.17
2788	X72688	L.stagnalis LS-rab1 mRNA	0.2
2789	XM_033154	Homo sapiens syntrophin, gamma 1 (SNTG1), mRNA	0.021
2790	AB017186	Clostridium perfringens genes of hem operon, complete cds	1.4
		Solanum allophyllum NADH dehydrogenase subunit (ndhF) gene,	
2791	U47416	chloroplast gene encoding chloroplast protein, partial cds	0.2
		Human small nuclear ribonucleoprotein (snRNP) E gene, exon 5 and	
2792	M21258	Alu repeats copy N and O	0.051
		Staphylococcus aureus dehydroquinate synthase (aroB) gene, 3' end	
		cds; 3-phosphoshikimate-1-carboxyvinyltransferase (aroA) gene,	
2793	L05004	complete cds; ORF3, complete cds	0.21
-		Uma scoparia isolate ROM3406 cytochrome b gene, partial cds;	
2794	AF302005	mitochondrial gene for mitochondrial product	2
2795	AF288209	Homo sapiens beta galactosyltransferase bGalT7 mRNA, complete cds	2
2796	AB048897	Macaca fascicularis brain cDNA, clone:QnpA-17253	1.7
2797	AK025054	Homo sapiens cDNA: FLJ21401 fis, clone COL03678	0.071
2798	U39886	Bos taurus Y-chromosome specific genomic sequence	0.002
2799	AJ277649	Caenorhabditis elegans mRNA for CHE-14 protein	0,6
		Xenopus laevis bone morphogenetic protein 4 (BMP-4) gene,	
2800	AF058764	complete cds	0.56
		Plasmodium falciparum chromosome 2, section 35 of 73 of the	
2801	AE001398	complete sequence	0.15
2802	AF079456	O'nyong-nyong virus strain SG650, complete genome	0.003
2803	AF087913	Human endogenous retrovirus HERV-P-T47D	6
2804	AB005053	Anomala cuprea entomopoxvirus gene for spheroidin, complete cds	1.8
2805	AF027656	Homo sapiens cholesteryl ester transfer protein gene, promoter region	0.004
2806	XM_031478	Homo sapiens vacuolar protein sorting protein 18 (VPS18), mRNA	1.7
2807	XM_001718	Homo sapiens KIAA0475 gene product (KIAA0475), mRNA	0.023
2808	L21906	Human troponin I, slow-twitch isoform (TNNI1) gene, exons 2-4	1.5
		Homo sapiens IQ motif containing GTPase activating protein 2	
2809	NM_006633	(IQGAP2), mRNA	5E-39
		Helicobacter pylori, strain J99 section 102 of 132 of the complete	
2810	AE001541	genome	0.2
2811	M63577	S.cerevisiae SFP1 gene, complete cds	0.023
2812	XM_004980	Homo sapiens cystic fibrosis transmembrane conductance regulator, ATP-binding cassette (sub-family C, member 7) (CFTR), mRNA	0.013
2813	AJ305053	Caprine arthritis-encephalitis virus proviral partial pol gene for Pol protein (reverse transcriptase), isolate CAEV Br/UFRGS-2/C767	0.5

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Mus musculus 18 days pregnant adult female placenta and extra	
		embryonic tissue cDNA, RIKEN full-length enriched library,	
2814	AK014450	clone:3830422A13, full insert sequence	0.075
2815	X99691	B.taurus DNA for agouti gene	2
		H.sapiens CpG island DNA genomic Mse1 fragment, clone 152h5,	
2816	Z64630	forward read cpg152h5.ft1a	0.14
2817	AE257071	Ovine pulmonary adenocarcinoma virus, complete sequence; and	
2017	AF357971	flanking Ovis aries sequence	4.9
2818	AE003336	Drosophila melanogaster genomic scaffold 142000013385800, complete sequence	
2819	U72236	Dictyostelium discoideum ModA (modA) gene, complete cds	0.16
2019	072230	Homo sapiens gap junction protein, beta 3, 31kD (connexin 31)	0.61
2820	XM_018110	(GJB3), mRNA	5.1
2020	22.12.010110	Capsicum annuum gene encoding 5-epi-aristolochene synthase, exons	3.1
2821	AJ005588	1 to 7	0.23
2822	AK022842	Homo sapiens cDNA FLJ12780 fis, clone NT2RP2001813	0.074
	1-4-022012	Treate supress estat 1 set 12/00 mg erone 1412/14 2001015	0.074
2823	AF138281	Arabidopsis thaliana phospholipase D-gamma-2 mRNA, complete cds	0.071
2824	AB040882	Homo sapiens mRNA for KIAA1449 protein, partial cds	0.18
		Mus musculus Cctb gene for chaperonin containing TCP-1 beta	0,10
2825	AB022156	subunit, complete cds	0.5
		Mus musculus, RIKEN cDNA 1200007D18 gene, clone MGC:7601	
2826	BC005516	IMAGE:3494155, mRNA, complete cds	0.51
2827	AK025207	Homo sapiens cDNA: FLJ21554 fis, clone COL06330	4.4
2828	NM_018798	Mus musculus ubiquilin 2 (Ubqln2), mRNA	0.043
		H. sapiens (D17S842) DNA segment containing (CA) repeat; clone	
2829	Z23770	AFM240xe5; single read	0.00004
		Mus musculus, Similar to proteasome (prosome, macropain) subunit,	
		alpha type 1, clone MGC:6546 IMAGE:2655483, mRNA, complete	
2830	BC005762	cds	0,67
2831	AB047932	Macaca fascicularis brain cDNA, clone:QnpA-10158	0.68
2832	U19765	Human nucleic acid binding protein gene, complete cds	0.18
2833	AF200327	Plasmodium chabaudi unknown genes	0.028
2834	U67544	Methanococcus januaschii section 86 of 150 of the complete genome	0.14
		Homo sapiens, Similar to hypothetical protein, clone MGC:13367	
2835	BC010357	IMAGE:4249317, mRNA, complete cds	2E-55
2836	AE007471	Streptococcus pneumoniae section 154 of 194 of the complete genome	5,2
2837	AJ320185	Solanum tuberosum sn1 gene for snakin-1, exons 1-2	0.44
		, A. A. A. A. A. A. A. A. A. A. A. A. A.	
2838	AF039954	Homo sapiens CC chemokine LCC-1 precursor, gene, complete cds	0.002
2839	Y13631	Clostridium botulinum P-21, P-47 ntnh, bonT genes	0.14
		Homo sapiens mRNA; cDNA DKFZp547J125 (from clone	
2840	AL390181	DKFZp547J125)	0.019

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALU
		Diadema antillarum isolate dmid3 tRNA-Lys gene, partial sequence;	
		ATPase 8 gene, complete cds; and ATPase 6 gene, partial cds;	1
2841	AF366200	mitochondrial genes for mitochondrial products	6.1
2842	AJ237688	Canis familiaris polymorphic marker clone XA07	0.53
		Mus musculus galectin-11 mRNA, variant b, complete cds,	
2843	AF244979	alternatively spliced	0.25
		Homo sapiens mRNA; cDNA DKFZp547J184 (from clone	
2844	AL390174	DKFZp547J184)	0.21
		Human clotting factor VIII gene, junction regions of the deletion of	
2845	U80230	exons 15 through 20 and LINE-1-like repeat region	2E-09
		Gallus gallus neuron-glia adhesion molecule (Ng-CAM) gene, partial	
2846	U31086	cds, exons 1 and 2	0.064
2847	D16685	Rice gene for lactate dehydrogenase, complete cds	0.003
2848	AB007459	Chara corallina ccm1 mRNA for myosin, complete cds	0.067
2849	U63328	Pissodes strobi vitellogenin (Vtg) mRNA, partial eds	0.68
2850	U50727	Human p40-phox (NCF4) gene, exon 8	0.023
2851	NM_017190	Rattus norvegicus Myelin-associated glycoprotein (Mag), mRNA	6.5
2852	AF361105	Homo sapiens interleukin 9 (IL9) gene, complete cds	0.37
		Homo sapiens genomic DNA, chromosome 21q22.2, clone:D34.	
2853	AP001416	LB7T-ERG region, complete sequence	1.5
2854	Y16771	Phylica pubescens chloroplast partial trnL gene & intergenic spacer	0.6
		Escherichia coli O157:H7 EDL933 genome, contig 3 of 3, section 243	
2855	AE005624	of 290	0.075
		Microbotryum violaceum ammonium transporter MEPa (MEPa)	
2856	AF159568	mRNA, complete cds	0.7
2857	M34149	S.cerevisiae mitochondrial gap 12 sequence	0.51
2858	D16541	Chicken DNA for connectin(titin), partial sequence	0.064
2859	AC007039	Homo sapiens BAC clone RP11-263A15 from Y, complete sequence	e-149
2860	AJ294714	Pichia anomala leu2 gene for beta-isopropylmalate dehydrogenase	0.056
2861	XM_051160	Homo sapiens KIAA1463 protein (KIAA1463), mRNA	0.22
2862	XM_031527	Homo sapiens leucine-rich PPR-motif containing (LRPPRC), mRNA	0.0003
		Plasmodium yoelii yoelii clone 11 235 kDa rhoptry protein (IIa.2)	
2863	AF323443	gene, partial cds	0.02
		Staphylococcus aureus partial ORF292 and ORF271, ORF331 and	
2864	Y14324	ORF314	0.62
2865	BC007568	Homo sapiens, clone IMAGE:3028427, mRNA, partial cds	0.16
2866	U67518	Methanococcus jannaschii section 60 of 150 of the complete genome	0.67
2867	AJ133742	Sus scrofa ryr3 gene (partial), exon 2	0.68
2868	AF116520	Mus musculus periplakin (ppl) gene, exons 2-14	0.083
2869	D38044	Human gene for Ah-receptor, exon 7-9	0.25

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID	A COTTON!	222000	
NO	ACCESSN	DESCRIP	P VALUE
2870	XM_010162	Homo sapiens methyl CpG binding protein 2 (Rett syndrome) (MECP2), mRNA	1.7
		Lavatera phoenicea isolate 804 internal transcribed spacer 1, partial	
		sequence, 5.8S ribosomal RNA, complete sequence and internal	1
2871	AF303030	transcribed spacer 2, partial sequence	0.01
2872	AE002274	Chlamydia muridarum, section 6 of 85 of the complete genome	1.8
		Homo sapiens p53DINP1 gene for p53DINP1a, p53DINP1b, complete	İ
2873	AB062056	cds, alternative splicing	8E-26
		Bacillus anthracis plasmid pXO1 resolvase X (resX) and transposase	
2874	AF150965	X (traX) genes, complete cds	1.6
		Human chromosome 16 duplicated adrenoleukodystrophy (ALD)	1
2875	U90292	gene, 5' breakpoint sequence	3E-41
2876	U75361	Rattus norvegicus Munc13-3 mRNA, complete cds	3E-18
2877	AF006686	Plecotus auritus clone Paur06 microsatellite sequence	5E-10
			1
2878	NM_009694	Mus musculus apolipoprotein B editing complex 2 (Apobec2), mRNA	0.00009
2879	AF315316	Vigna radiata ACC oxidase gene, complete cds	0.003
2880	AE005714	Caulobacter crescentus section 40 of 359 of the complete genome	1.5
		Human DNA sequence from clone RP11-45G20 on chromosome 6,	
2881	AL451135	complete sequence [Homo sapiens]	1.9
2882	AF315316	Vigna radiata ACC oxidase gene, complete cds	0,003
2883	NM_009694	Mus musculus apolipoprotein B editing complex 2 (Apobec2), mRNA	0,00008
2884	AB047932	Macaca fascicularis brain cDNA, clone:QnpA-10158	0.16
2885	U18671	Human Stat2 gene, complete cds	0.23
		Bombyx mori copg2 mRNA for nonclathrin coat protein gamma2-	
2886	AB040670	COP, partial cds	0.074
		Mus musculus adult male thymus cDNA, RIKEN full-length enriched	
2887	AK017987	library, clone:5830443J22, full insert sequence	0.035
		Plasmodium falciparum chromosome 2, section 56 of 73 of the	
2889	AE001419	complete sequence	0.12
1		Eriosoma lanigerum 12S small subunit ribosomal RNA gene, partial	1
Į l		sequence; tRNA-Val gene, complete sequence; and 16S large subunit	
		ribosomal RNA gene, partial sequence; mitochondrial genes for	
2890	AF275225	mitochondrial products	0.35
2891	AF372971	Arabidopsis thaliana AT3g23580/MDB19_7 mRNA, complete cds	0.53
		Homo sapiens partial ARNT gene for aryl hydrocarbon receptor	
2892	AJ404851	nuclear translocator, exon 4	3.7
2893	AJ012380	Fugu rubripes partial rara gene, exons 4-10	1.4
2894	AB064593	Escherichia coli icdA gene for isocitrate dehydrogenase, complete cds	0.027
		Drosophila melanogaster disco-interacting protein (dip1) gene,	
		alternatively spliced products, complete cds; and TO42 gene, partial	
2895	AF182444	cds	1.6

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
2896	AY020760	Oryza sativa microsatellite MRG3085 containing (TA)X13, closest to marker R2976, genomic sequence	0.007
2005	177005010	Lactococcus lactis subsp. lactis IL1403 section 80 of 218 of the	
2897 2898	AE006318 X01973	complete genome	0.53
2098	A01973	Mouse gene for interferon alpha 4 (Mu IFN-alpha 4) Candida albicans gamma-glutamylcysteine synthetase (GCS1) gene.	3.9
2899	AF176677	candida aiorcans gamma-giinamyicysteine synthetase (GCS1) gene,	0.07
2900	AF109009	Prolaupala kukui clone 1092 transposon Lau1 sequence	3.6
2901	D49739	Chicken mRNA for leucine zipper protein, complete cds	2.4
2902	U34610	Mus musculus alpha-1(XVIII) collagen (COL18A1) gene, exon 6 and 7	0,006
2903	AK026763	Homo sapiens cDNA: FLJ23110 fis, clone LNG07802	0,34
2904	AF343914	Campylobacter jejuni NCTC 11828 LOS biosynthesis cluster, partial sequence	0.024
2905	AE002722	Drosophila melanogaster genomic scaffold 142000013385665, complete sequence	0.007
2906	XM_027242	Homo sapiens KIAA0805 protein (KIAA0805), mRNA	1.1
2907	Y15082	Homo sapiens p14.5 gene, partial exon 1 and promoter; hPOP1 gene	0.65
2907	XM 009536	exon 1 and promoter and bidirectional promoter region Homo sapiens GTPase GES; REM protein (REM), mRNA	0.67
2908	YMT_005330	Homo sapiens G1Pase GES; REM protein (REM), mRNA	5.9
2909	XM 036351	Homo sapiens nuclear transcription factor Y, alpha (NFYA), mRNA	1E-32
2910	AB037851	Homo sapiens mRNA for KIAA1430 protein, partial cds	0.58
2911	XM 002943	Homo sapiens similar to PRO0327 protein (H. sapiens) (LOC92570), mRNA	2,4
2912	AF069179	Lysiphlebus cardui NADH dehydrogenase 1 gene, mitochondrial gene encoding mitochondrial protein, partial cds	0.008
2913	AJ243545	Arabidopsis thaliana gene for 67 kD chloroplastic RNA-binding protein, P67	0.54
2914	AJ295795	Isolepis producta chloroplast trnL gene intron	2.2
2915	AJ311050	Medicago sativa pr10.2 gene for class 10 PR protein	0.26
2916	AK021512	Homo sapiens cDNA FLJ11450 fis, clone HEMBA1001432	0.25
2917	AK002979	Mus musculus adult male brain cDNA, RIKEN full-length enriched library, clone:0710001P07, full insert sequence	0.023
		Streptococcus mitis phage SM1 PbIA (pbIA) and PbIB (pbIB) genes,	
2918	AY007505	complete cds; lysin (lys) gene, partial cds; and unknown genes	0.026
2919	AK022043	Homo sapiens cDNA FLJ11981 fis, clone HEMBB1001317	0.69
2920	AF162969	Mycoplasma capricohm capricolum strain 960038 fructose biphosphate aldolase gene, partial cds	0.064
2921	XM 051364	Homo sapiens similar to immediate early protein (H. sapiens) (LOC93434), mRNA	0.086
2922	XM 027642	Homo sapiens KIAA0349 protein (KIAA0349), mRNA	0.24
2923	XM 045819	Homo sapiens cofilin 2 (muscle) (CFL2), mRNA	1.9
2924	AF146362	Drosophila melanogaster Huntington disease protein homolog (Hsap\HD) mRNA, complete cds	0.23

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
2925	U91993	Solanum chacoense transposable element Sch2, sequence	0.027
2926	AK021782	Homo sapiens cDNA FLJ11720 fis, clone HEMBA1005293	2.3
		Carsonella ruddii ribosomal protein L11 (rpL11), ribosomal protein L1 (rpL1), ribosomal protein L10 (rpL10), ribosomal protein L7/L12	
2927	AF274444	(rpL7/L12), beta subunit of RNA polymerase (rpoB), beta-prime subunit of RNA polymerase (rpoC), ribosomal protein S12 >	2
2721	AU2/4444	Simian immunodeficiency virus isolate P2S1 patient P2, gp120 (env)	
2928	AF003047	gene, partial cds	0.064
2929	AK022102	Homo sapiens cDNA FLJ12040 fis, clone HEMBB1001944	0.064
2929	AK022102	Homo sapiens BAC clone CTB-34F8 from 7p15-p21, complete	U
2930	AC004239	sequence	. 0.001
2930	AC004239		. 0.001
2931	47104010	Strumaria truncata trnL gene, partial sequence; chloroplast gene for	
2931	AF104819	chloroplast product	0.25
2022	AT151016	Trimeresurus flavomaculatus cytochrome b (cytb) gene, partial cds;	
2932	AF171916	mitochondrial gene for mitochondrial product	0.21
2933	AJ011596	Homo sapiens trapped 3' terminal exon, clone B2E8	0.0003
2934	AF269137	Homo sapiens GABAA receptor gamma 3 subunit (GABRG3) gene, exon 3	0.8
255.	14 207157	Arabidopsis thaliana AATL1 gene for amino acid transporter-like	0.0
2935	AB030586	protein 1, complete cds	1.9
		Cronartium appalachianum isolate Ca-1 18S ribosomal RNA gene,	
		partial sequence; internal transcribed spacer 1, 5.8S ribosomal RNA	
		gene and internal transcribed spacer 2, complete sequence; and 26S	
2936	L76484	ribosomal RNA gene, partial sequence	0.71
2937	AF366570	Trypanosoma congolense calreticulin mRNA, complete cds	0.084
2938	NM_004682	Homo sapiens PC4 and SFRS1 interacting protein 1 (PSIP1), mRNA	2
		Homo sapiens mRNA; cDNA DKFZp586J1623 (from clone	1
2939	AL050219	DKFZp586J1623)	1.8
		Plasmodium falciparum knob-associated histidine-rich protein	
2940	AF275687	(KAHRP) gene, complete cds	0.009
		Vigna umbellata maturase-like protein (matK) gene, complete cds;	
2941	AF142701	chloroplast gene for chloroplast product	0.56
		Mitthyridium undulatum small ribosomal protein 4 (rps4) gene,	
2942	AF226775	partial cds; chloroplast gene for chloroplast product	0.073
		Homo sapiens similar to hypothetical protein FLJ21463 (H. sapiens)	
2943	XM_034735	(LOC90883), mRNA	0
		Mus musculus adult male thymus cDNA, RIKEN full-length enriched	
2944	AK017899	library, clone:5830405M20, full insert sequence	0.28
2945	Y10159	D.discoideum racGAP gene	6.2
		Homo sapiens hypothetical gene supported by NM_000181	
2946	XM_011562	(LOC92707), mRNA	0.025
2947	AF191257	Trichuris trichiura microsatellite ttstr14	0,078
2948	Y08449	E.caballus DNA segment containing CA microsatellite VHL137	0.00003

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Mus musculus 18 days embryo cDNA, RIKEN full-length enriched	
2949	AK003959	library, clone:1110029G12, full insert sequence	0.077
2950	NM_012779	Rattus norvegicus Aquaporin 5 (Aqp5), mRNA	2.2
2951	AJ404295	Borelis schlumbergeri 18S rRNA gene, isolate 191	0.25
		Plasmodium falciparum chromosome 2, section 2 of 73 of the	l
2952	AE001365	complete sequence	0.49
2954	Y13434	Ovis aries mRNA for thyroid stimulating hormone receptor	1.8
		Homo sapiens SDHD gene for small subunit of cytochrome b of	l
2955	AB026906	succinate dehydrogenase, complete cds	0.001
		Homo sapiens, hypothetical protein FLJ20534, clone MGC:13432	1
2956	BC010367	IMAGE:4334172, mRNA, complete cds	4
2957	AJ251957	Ciona intestinalis mRNA for nuclear lamin (lamin L2 gene)	2
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
2958	AK016695	library, clone:4933406J08, full insert sequence	2E-14
2959	Z13985	M.musculus control region sequence DNA	0.072
		Human DNA sequence from clone CTD-2053G21 on chromosome	
·2960	AL591590	Xq13.3-21.2, complete sequence [Homo sapiens]	0.007
2961	AB002807	Glycine max DNA for nodulin 35, partial cds	0.24
		Plasmodium falciparum chromosome 2, section 36 of 73 of the	
2962	AE001399	complete sequence	0.003
2963	AF298224	Homo sapiens RPCI-II 289E13 CD21 gene, partial cds	0.00001
		Homo sapiens similar to hypothetical protein PRO2822 (H. sapiens)	
2964	XM_018458	(LOC93537), mRNA	0.21
2965	AF288778	Homo sapiens erythroleukemia translocation breakpoint region	0.068
		Drosophila melanogaster gamma-tubulin (GTUB) gene, partial cds:	
		and recombination repair protein 1 (RRP1) and thymidylate synthase	i
		(TS) genes, complete cds; and RNA-binding protein (RBP9) gene,	l
2966	AF073994	partial sequence	0.14
		Human DNA sequence from clone 346P11 on chromosome Xq21,2-	
2967	AL121822	21.33, complete sequence [Homo sapiens]	0.0000001
		Mesotaenium caldariorum clone mesphy1b phytochrome gene.	
2968	U31284	complete cds	0.59
2969	XM_008806	Homo sapiens mitogen-activated protein kinase 4 (MAPK4), mRNA	1.8
2970		Homo sapiens KIAA1209 protein (KIAA1209), mRNA	0.001
2971	AF179711	Hylobates sp. RH50 glycoprotein (RHAG) gene, intron 9	0.56
		Mus musculus, apoptosis inhibitory protein 5, clone MGC:5830	
2972	BC007133	IMAGE:3594358, mRNA, complete cds	5.4
2973	AF282018	Homo sapiens clone 15qtel ctg0082 c262bt3 c290bt3 sequence	2E-53
		Pseudomonas aeruginosa PA01, section 351 of 529 of the complete	
2974	AE004790	genome	1.8
		Homo sapiens similar to seven transmembrane domain orphan	
2975	XM 003050	receptor (H. sapiens) (LOC63108), mRNA	0.27
		Mus musculus myo-inositol monophosphatase 1 (Impa1) gene, exons	L
2976	AF353728	1 through 7	1.2
22.00	505720	1	

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Mus musculus adult female placenta cDNA, RIKEN full-length	
2977	AK005535	enriched library, clone:1600025H15, full insert sequence	0.55
2978	AJ002479	Medicago truncatula ENBP1 gene, exons 1 to 12	0.64
		Homo sapiens Ras-like GTP-binding protein (RAB27A) gene, exons	
2979	AF154836	1b and 2	0.000001
2981	AF260819	Plasmodium falciparum CG1 protein gene, complete cds	0.73
2982	AE006714	Sulfolobus solfataricus section 73 of 272 of the complete genome	0.02
2984	NM_004114	Homo sapiens fibroblast growth factor 13 (FGF13), mRNA	6.7
		Buthus martensii putative potassium ion channel blocker TXKs1	
2985	AF155369	mRNA, complete cds	0.72
		Homo sapiens similar to MHC class I region ORF (H. sapiens)	
2986	XM_041836	(LOC91955), mRNA	0.18
		Microcephala discoidea NADH dehydrogenase (ndhF) gene, partial	
2987	AF153668	cds; chloroplast gene for chloroplast product	1.7
		Drosophila buzzatii alpha-esterase 3 (aE3) and alpha-esterase 4b	
2988	AF216211	(aE4b) genes, complete cds	0.073
		Dermatophagoides farinae paramyosin-like allergen mRNA, partial	
2989	AF352244	cds	0.025
2990	XM_007409	Homo sapiens alkylation repair; alkB homolog (ABH), mRNA	0.21
2991	AF005779	Sus scrofa neuropeptide Y Y1 receptor mRNA, complete cds	0.66
		Arabidopsis thaliana hydrophobic protein RCI2B (RCI2B) and	
2992	AF264749	hydrophobic protein RCI2A (RCI2A) genes, complete cds	0.022
		Homo sapiens mRNA; cDNA DKFZp434F1872 (from clone	
2994	AL162033	DKFZp434F1872)	0.69
		Mus musculus adult male hippocampus cDNA, RIKEN full-length	1 _
2995	AK013593	enriched library, clone:2900024O09, full insert sequence	0.7
2006		Plasmodium falciparum clone 3D7 unconventional myosin PfM-B	
2996	AF222716	gene, complete cds	0.027
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
2997	AK006599	library, clone:1700034J06, full insert sequence	2.2
2998	AF010509 AF070566	Plasmodium falciparum microsatellite TA39 sequence	0.001
		Homo sapiens clones 24611 and 24705 mRNA sequence	0.000003
3000	AF181676	Homo sapiens clone 204112 polymorphic microsatellite sequence	0.074
2001	3.6151.60	Chinese hamster glucose-regulated protein GRP78 mRNA, complete	
3001	M17169	cds	0.008
3002	NM_031236	Rattus norvegicus alpha 1,2-fucosyltransferase (Fut1), mRNA	0.69
3003	V2.6 016221	Homo sapiens region containing TBX3-iso protein; T-box 3 (ulnar	0.50
3003	XM_016321	mammary syndrome) (LOC82353), mRNA	0.73
3004	DC005250	Homo sapiens, Similar to plasminogen, clone MGC:12496	0.00000
3004	BC005379 AF276176	IMAGE:3934860, mRNA, complete cds	0.000001
3005		Scaphirhynchus platorynchus microsatellite Spl-107 sequence	0.009
3006	XM_010048 NC 001807	Homo sapiens like mouse brain protein E46 (E46L), mRNA	0.47
3007	NC_00180/	Human mitochondrion, complete genome	1E-10
3008	AE006103	Pasteurella multocida PM70 section 70 of 204 of the complete genome	0.4

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Lactococcus lactis subsp. lactis IL1403 section 28 of 218 of the	
3009	AE006266	complete genome	0.2
		Dehalospirillum multivorans GTP cyclohydrolase II (ribA) gene,	
		partial cds; and 3,4-dihydroxy-2-butanone-4-phosphate synthase	
		(ribB), riboflavin synthase (ribH), tetrachloroethene reductive	
3010	AF022812	dehalogenase catalytically active subunit (pceA), and tetrachl>	0.21
3011	AJ243901	Mycoplasma hominis p75 gene, strain PG21	1.4
		Homo sapiens hypothetical protein DKFZp761C121	
3013	XM_027894	(DKFZp761C121), mRNA	5E-33
3014	AJ390525	Candida albicans ORF DNA, clone 64gB	0.52
		Buchnera aphidicola plasmid pBUs1, repA2 gene, leuA gene and	
3015	AJ006873	ORF1	0.013
3016	L07898	Entamoeba histolytica alpha-1 tubulin (tub1) gene, complete cds	0.024
3017	L25107	Human LIS mRNA	1.2
		Homo sapiens mRNA; cDNA DKFZp434L098 (from clone	
3018	AL122122	DKFZp434L098)	0.007
3019	V01270	Rattus norvegicus genes for 18S, 5.8S, and 28S ribosomal RNAs	2E-10
3020	AK023061	Homo sapiens cDNA FLJ12999 fis, clone NT2RP3000324	0.12
3021	U80449	Caenorhabditis elegans cosmid R10A10	0.29
3022	AF355473	Oryzias latipes germ cell-less protein (gcl) mRNA, complete cds	2.3
		Homo sapiens hypothetical gene supported by AL117650	
3023	XM_037811	(LOC91349), mRNA	0.39
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
3024	AK016492	library, clone:4931430N09, full insert sequence	0.004
		Homo sapiens similar to DKFZP564K247 protein (H. sapiens)	
3025	XM_011189	(LOC91537), mRNA	4
		Mus musculus, Similar to ribosomal protein L19, clone MGC:6500	
3026	BC010710	IMAGE:2648593, mRNA, complete cds	5.1
3027	XM_017923	Homo sapiens hypothetical protein FLJ12660 (FLJ12660), mRNA	4.1
3028	AC016671	Homo sapiens BAC clone RP11-12J24 from Y, complete sequence	0.012
		Campylobacter jejuni major cell-binding factor (peb1) and receptor	
3029	L13662	genes, complete cds	0.044
		M.musculus pF2 genomic DNA, homology with mouse repetitive	
3030	X77172	elements B1, B2 and human Alu	0.008
3031	AF074841	Campylobacter lari tRNA-Ala and tRNA-Ile genes, complete sequence	0.016
		Homo sapiens Fanconi anemia complementation group D2 protein	
3032	AF272846	(FANCD2) gene, exon 1	0.065
		Drosophila americana texana G-3-P dehydrogenase (alphaGpdh)	
3033	D50090	gene, exon1-8, complete cds	0.13
		Plasmodium falciparum chromosome 2, section 54 of 73 of the	
3034	AE001417	complete sequence	0.15
		Homo sapiens cDNA FLJ11145 fis, clone PLACE1006626, highly	
3035	AK002007	similar to Homo sapiens mRNA for KIAA0928 protein	0.026

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			r
NO	ACCESSN	DESCRIP	P VALUE
		Human DNA sequence from clone RP11-245H22 on chromosome 6	1
3036	AL512432	Contains GSSs, complete sequence [Homo sapiens]	0.00003
2027	17070100	Glycine max SG-05 gene for thiamin biosynthetic enzyme, complete	
3037	AB030490	cds	0.003
		Saccharomyces cerevisiae DA-box protein Smc2p (SMC2) gene,	l
3038	U05820	complete cds, and assimilatory sulfite reductase (MET10) gene, partial cds	
3039	AF098761	Gracilaria gracilis microsatellite Gg155R sequence	0.07
3037	AF038701	Carassius auratus clone gf-40 glutamic acid decarboxylase isoform 67	0.07
3041	AF045595	(GAD67) mRNA, complete cds	2
3042	AJ404308	Sorites marginalis 18S rRNA gene, isolate 836	0.009
	14101500	Streptococcus pyogenes M1 GAS strain SF370, section 24 of 167 of	0.007
3043	AE006495	the complete genome	6
			<u>`</u>
3044	XM 030781	Homo sapiens enigma (LIM domain protein) (ENIGMA), mRNA	3E-18
3045	AC024748	Caenorhabditis elegans cosmid Y110A2AR, complete sequence	0.15
		Human DNA sequence from clone XXvac-BA141F18 on chromosome	
3046	AL512282	X, complete sequence [Homo sapiens]	0.06
		Otolemur garnetti color vision opsin gene, exons 5 and 6, and	
3047	AF009980	complete cds	0.17
		Carex exsiccata tRNA-Thr, partial sequence; trnT-trnL intergenic	
1		spacer, tRNA-Leu, and trnL-trnF intergenic spacer, complete	
		sequence; and tRNA-Phe, partial sequence; chloroplast genes for	
3048	AF284953	chloroplast products	0.23
3049	AE004036	Xylella fastidiosa 9a5c, section 182 of 229 of the complete genome	4.3
3050	X94767	H.sapiens RP3 gene (XLRP gene 2)	0.002
3030	A34707	Dictyostelium discoideum spore coat structural protein SP65 (cotE)	0.002
3051	AF279135	gene, complete cds	0.003
3052	AF277452	Rattus norvegicus Pkd1 mRNA, partial cds	0.12
3053	AF327114	Homo sapiens clone 8qtel c19t7 sequence	8E-17
3054	AB052201	Macaca fascicularis brain cDNA, clone:QnpA-20394	0.59
3055	AJ235314	Antilocapra americana mitochondrial DNA for D-loop	0.17
		Homo sapiens mitochondrial GTP-binding protein 1 (GTPBP3) gene,	
3056	AF361481	complete cds; nuclear gene for mitochondrial product	0.009
3057	AE006879	Sulfolobus solfataricus section 238 of 272 of the complete genome	1.6
2050		Legionella pneumophila ORF X (partial), recA gene, recX gene and	
3058	AJ277756	sya gene (partial)	0.21
3059	U73044	Schizosaccharomyces pombe ARS binding protein 2 (abp2) mRNA, complete cds	0.050
3039	073044	witipiete dis	0.058
3060	AF043123	Homo sapiens interleukin-7 receptor precursor (IL7R) gene, exon 1	0.2
2000	.11 073123	Atomo suprems ametrement-7 receptor precursor (11.77c) gene, exon 1	0.2
3061	L43603	Gallus gallus Na+/K+-ATPase alpha-1 subunit gene, promoter region	0.48
		- F Paradian Bone, Promotor region	0.10

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		The state of the s	T
NO	ACCESSN	DESCRIP	P VAL
3062	X68976	N.viridescens mRNA for Hoxd-10	0.09
		Helianthus annuus Dc3 promoter-binding factor-2 (DPBF-2) mRNA.	1
3063	AF001454	complete cds	0.1
3064	AF086062	Homo sapiens full length insert cDNA clone YZ06B11	3E-2
3065	U67234	Human clone HS5.29 Alu-Ya5 sequence	0.08
3066	AF030199	Mus musculus type 1 sigma receptor gene, complete cds	0.4
		Plasmodium falciparum glucosephosphate isomerase (ggi.R1) gene,	
3067	J05544	complete cds	0.2
3068	X17455	Rat Bmyc gene 5'-fragment	0.00
3069	AF298207	Dictyostelium discoideum transposon thug-S, complete sequence	0.00
5007	111230207	Sitty contrain discondenti transposori unig 5, complete sequence	0.00
3070	NM 030238	Mus musculus dynein, cytoplasmic, heavy chain 1 (Dnchc1), mRNA	1.7
3070	144_050250	Homo sapiens BABP gene for bile acid-binding protein [AKR 1C2].	1.7
3071-	AB032151	exon 1 to 7	0.2
3072	NC 001481	Feline calicivirus, complete genome	1.9
3072	NC_001481		1.3
3073	U49266	Mus musculus growth hormone receptor/binding protein gene, exons	0.1
3073	049200	9-10, and partial cds	0.1
3074	Z24279	H. sapiens (D1S489) DNA segment containing (CA) repeat; clone	١.,
3074	Z24219	AFM309ve9; single read	0.1
		Rhagoletis conversa NADH-dehydrogenase subunit 4 (ND4) gene,	
		partial sequence; NADH-dehydrogenase subunit 4L (ND4L) gene,	
2055	4 T2200 T2 4	complete sequence; and tRNA-Thr gene, partial sequence;	١.,
3075	AF228724	mitochondrial genes for mitochondrial products	0.04
		Oryza longistaminata receptor kinase-like protein (Xa21) gene,	
3076	U72723	complete cds and family member C, pseudogene	0.2
3077	U55184	Human G protein Golf alpha gene, exon 12 and complete cds	1E-
		Mus musculus clone UWGC:mbac10gap from 14D1-D2 (T-Cell	
3078	AC006119	Receptor Alpha Locus), complete sequence	0.000
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
3079	AK015900	library, clone:4930526F13, full insert sequence	0.7
		Homo sapiens mRNA; cDNA DKFZp434J1618 (from clone	
3080	AL133637	DKFZp434J1618); partial cds	0.07
3081	S74063	c-raf=raf protein [Xenopus laevis, mRNA, 2736 nt]	0.5
3082	NM_014498	Homo sapiens type II Golgi membrane protein (GPP130), mRNA	1.9
3083	AF238234	Entamoeba histolytica diaphanous protein (dia) gene, partial cds	2.1
		Homo sapiens SWI/SNF related, matrix associated, actin dependent	
3084	XM_010729	regulator of chromatin, subfamily f, member 1 (SMARCF1), mRNA	0.6
		Mus musculus 10 day old male pancreas cDNA, RIKEN full-length	
3085	AK007542	enriched library, clone:1810019D05, full insert sequence	0.6
		Human T-cell lymphotropic virus type 1 isolate ATLL long terminal	
3086	AF228977	repeat, partial sequence	0.00
3087	AC091698	Homo sapiens clone RP11-596J6, complete sequence	0.0
3088	AF233882	Homo sapiens plakoglobin (JUP) gene, exons 1 through 7	4E-8

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Drosophila koepferae KO4 San Luis, Argentina, Gandalf transposable	
3089	U29468	element DNA< partial sequence	0.2
		Mycoplasma hyopneumoniae strain ISU 232 YX1 (yx1) gene, partial	
		cds; p110 membrane protein precursor, gene, complete cds; and YX2	
3090	AF279292	(yx2) gene, partial cds	0.2
3091	AF391284	Homo sapiens 11p15.5 clone LOH11A, partial sequence	0.007
3092	AF259737	Ovis aries microsatellite MNS-64 sequence	0.66
3093	NM_030261	Mus musculus hypothetical protein MGC7182 (MGC7182), mRNA	6E-36
3094	AB047879	Macaca fascicularis brain cDNA, clone:QnpA-14270	0.00003
3095	X65607	H.sapiens MT1X gene for metallothionein 1X	0.023
		Pseudosymblepharis schimperiana tRNA-Leu, partial sequence; trnL-	
3096	AE221164	trnF intergenic spacer, complete sequence; and tRNA-Phe, partial	
3096	AF231164	sequence; chloroplast genes for chloroplast products	0.06
3097	AF290431	Mus musculus sequence downstream of Hyal1; and Fus2 gene, 5' UTR and exon 1	
3097	X96762	C.elegans mRNA for DNA topoisomerase I	0.028
3098	A30702	C. ciegans iniciva for DIVA topoisomerase i	0.028
		Salganea nigrita mitochondrial COII gene for cytochrome oxidase II,	
3099	AB036229	partial cds, collection_site:Lake Danao, Leyte, Philippines	0.53
5077	145050227	Staphylococcus aureus partial coa gene for coagulase, type F, isolate	0.55
3100	AJ309183	rs59	0.07
		Homo sapiens glutathione-S-transferase GSTT1 deletion/junction	0.07
3101	AF240785	region sequence	0.000004
		Homo sapiens synaptosomal-associated protein, 23kD (SNAP23).	
3102	XM_031737	mRNA (0.0000004
		Arabidopsis thaliana putative proline-rich protein (F23H24.15)	
3103	AY035097	mRNA, complete cds	0.18
		Anopheles farauti 3 12S small subunit ribosomal RNA gene, partial	
3104	AF121069	sequence; mitochondrial gene for mitochondrial product	0.065
3105	NM_016701	Mus musculus nestin (Nes), mRNA	0.23
		Homo sapiens retinoblastoma susceptibility protein (RB1) gene, exon	
3106	L41893	5, bases 44452-44939 in L11910	0.008
3107	Z78623	H. sapiens flow-sorted chromosome 6 TaqI fragment, SC6pA12B7	2E-21
2100	VA 6 0112/7	Homo sapiens NADH dehydrogenase (ubiquinone) 1 beta subcomplex,	27.24
3108	XM_041367	4 (15kD, B15) (NDUFB4), mRNA	2E-24
3109	AL117617	Homo sapiens mRNA; cDNA DKFZp564H0764 (from clone DKFZp564H0764)	0.17
3110	AK023907	Homo sapiens cDNA FLJ13845 fis, clone THYRO1000815	7E-42
2110	1 33023701	TIOMS SUPERING SEPTEM 12045 Ha, MORE 1111 NO 1000015	715-42
3111	M95396	Yeast putative transcriptional activator (ADA2) gene, complete cds	3.8
3112	NC 002670	Bacteriophage bIL311, complete genome	0.073
3113	XM_037870	Homo sapiens desmoplakin (DPI, DPII) (DSP), mRNA	2.2
		Homo sapiens hypothetical protein DKFZp547E052	
3114	NM 032276	(DKFZp547E052), mRNA	0.43

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
3115	AB048877	Macaca fascicularis brain cDNA, clone:QupA-16414	3E-30
		Mus musculus 13 days embryo head cDNA, RIKEN full-length	
3116	AK014209	enriched library, clone:3110053B16, full insert sequence	0.029
		Homo sapiens rhabdoid tumor deletion region protein 1 (RTDR1),	
3117	XM_037053	mRNA	9E-47
		Cinara pseudotaxifolia cytochrome oxidase II gene, partial cds;	
3118	AF156197	mitochondrial gene for mitochondrial product	0.18
3119	AK022006	Homo sapiens cDNA FLJ11944 fis, clone HEMBB1000684	0.003
3120	U23175	Caenorhabditis elegans cosmid F57F10, complete sequence .	0.022
		Candidatus Carsonella ruddii natural-host Arytaina genistae ATP	
		synthase alpha subunit (atpA) gene, partial cds; ATP synthase gamma	
3121	AF267198	subunit (atpG) gene, complete cds; and ATP synthase beta subunit (atpD) gene, partial cds	
3121	AF20/198	Homo sapiens CUB domain containing protein 1 (CDCP1) mRNA.	0.019
3122	AY026461	complete cds	0,5
3123	NC 002471	Typhlonectes natans mitochondrion, complete genome	0.45
3125	AF328531	Homo sapiens clone 9qtel c10t7 sequence	0.00001
5125	111 020001	Homo sapiens clone 11 pur alpha-associated ribosomal RNA gene,	0.00001
3126	AF397158	partial sequence	2E-37
3127	AF110970	HIV-1 isolate C-96BW11.06 country Botswana, complete genome	5.3
3128	AJ237690	Canis familiaris polymorphic marker clone XC08	1.3
3129	AE001176	Borrelia burgdorferi (section 62 of 70) of the complete genome	0.084
3130	AJ271333	Campylobacter hyoilei ORF1, ORF2, ORF3 and ORF4	6.7
		Merluccius productus Mpro-1 mitochondrial control region, partial	
3131	AF112247	sequence	0,0008
		Homo sapiens uncharacterized hypothalamus protein HT010 (HT010),	
3132	XM_002378	mRNA	4E-28
3133	AJ242726	Xenopus laevis mRNA for p21 activated kinase 2 (PAK2 gene)	.0,25
		Petunia x hybrida pollen-specific 1-aminocyclopropane-1-carboxylate	
3134	AF049711	synthase (ACS2) gene, complete cds	0.24
		Homo sapiens KIAA0001 gene product; putative G-protein-coupled receptor; G protein coupled receptor for UDP-glucose (KIAA0001).	
3135	XM 003091	mRNA	2.2
3136	Z33368	M.capricolum DNA for CONTIG MCABF	0.001
3130	233300	Plasmodium falciparum chromosome 2, section 70 of 73 of the	0.001
3137	AE001433	complete sequence	0.001
		Mus musculus adult male pituitary gland cDNA, RIKEN full-length	0.001
3138	AK017235	enriched library, clone;5330405D10, full insert sequence	0.7
		H.sapiens CpG island DNA genomic Msel fragment, clone 170g9,	
3139	Z57308	reverse read cpg170g9.rt1a	0.023
		Homo sapiens PAC clone RP5-998M2 from 7q33-q35, complete	
3140	AC005485	sequence	5.4
		Leishmania donovani ATPase 6 gene; kinetoplast gene for kinetoplast	

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		l same (same to same t	
NO	ACCESSN	DESCRIP	P VALUE
		Uncultured eubacterium plasmid pIE1130 DNA for repA gene, repC	
		gene, repB gene, mobA gene, mobB gene, mobC gene, cac gene, catIII	
3142	AJ271879	gene, sulII gene, strA gene, strB gene and aph(3')-I gene	0.26
3143	Y00165	Sus scrofa mRNA for link protein	0.079
		Caenorhabditis elegans transcription enhancer factor-1-like protein	
3144	AF283982	EGL-44 (egl-44) gene, complete cds	0.0007
	17001006	Pseudomonas aeruginosa PA01, section 487 of 529 of the complete	
3145	AE004926	genome	0.44
3146	17001700	Pseudomonas aeruginosa PA01, section 299 of 529 of the complete	
3146	AE004738	genome	1,4
3147	AC083824	Homo sapiens clone RPI1-91J18, complete sequence	0.59
3148	AY023127	Oryza sativa microsatellite MRG5452 containing (GCA)X9, closest to marker C402, genomic sequence	0.77
3140	A1023127	Homo sapiens similar to PRO0327 protein (H. sapiens) (LOC92570).	0.77
3149	XM 002943	mRNA	0.073
3147	7111_002545	Clostridium perfringens DNA for D-alanine; D-alanine ligase, cortical	0.073
3150	D88151	fragment-lytic enzyme	0.057
		Homo sapiens similar to tumor necrosis factor (ligand) superfamily,	0.037
3151	XM_009013	member 14 (H. sapiens) (LOC63472), mRNA	1.7
		Rattus norvegicus Carcinoembryonic antigen gene family (CGM1)	
3152	NM_012702	(Cgm1), mRNA	1.5
3153	AJ131041	Clogmia albipunctata mRNA for Hunchback protein, partial	0.019
3154	AE000720	Aquifex aeolicus section 52 of 109 of the complete genome	5.3
3155	X74789	Potato leaf roll virus RNA sequence	0.47
		Homo sapiens natural resistance-associated macrophage protein 2	
3156	AF064478	(NRAMP2) gene, exons 4 and 5	0.021
3157	M84761	Rat fatty acid synthase gene, complete cds	0.25
		Polycycnis ornata tRNA-Leu (trnL) gene, partial sequence; and trnL-F	
3158	AF239566	intergenic spacer region, complete sequence; chloroplast gene for chloroplast product	0.007
3136	AF 239300	Campylobacter jejuni NCTC 11828 LOS biosynthesis cluster, partial	0.007
3159	AF343914	sequence	0.025
3133	14 343714	Homo sapiens mRNA; cDNA DKFZp566G184 (from clone	0,023
3160	AL137443	DKFZp566G184)	0.99
3161	X54452	D. discoideum culmination spiA (Dd31) gene	0.71
3162	X80908	L.esculentum gene for fruit ripening polygalacturonase	0.069
		Homo sapiens nuclear factor of activated T-cells 5, tonicity-responsive	
3163	XM_007838	(NFAT5), mRNA	0.5
		Mus musculus KvLQT1 gene, KvLQT1-AS allele, partial intron	
3164	AF119385	sequence	0.17
3165	X86449	E.caballus DNA segment containing microsatellite	1E-25
3166	Z22800	H.sapiens microsatellite repeat	0.000004
3167	U22842	Lagos bat virus nucleoprotein gene, partial cds	0.027

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
1		Campylobacter jejuni PglB (pglB) gene, partial cds; and PglA (pglA),	
1		PglC (pglC), PglD (pglD), PglE (pglE), PglF (pglF), and PglG (pglG)	ļ
3168	AF108897	genes, complete cds	0.73
3169	AE000556	TV.P. I. decor	
3170	AK024295	Helicobacter pylori 26695 section 34 of 134 of the complete genome Homo sapiens cDNA FLJ14233 fis, clone NT2RP4000049	0.016
3170	AK024293	Mus musculus c-abl protein (c-abl) gene, exons 1 and 1B.	0.46
3171	U13835	alternatively spliced	4.5
3172	AJ132887	Homo sapiens telomere-junction DNA sequence, haplotype B	6E-13
	12102001	Homo sapiens hypothetical protein DKFZp761F2014	0E-13
3173	XM 007277	(DKFZp761F2014), mRNA	1.9
3174	L40470	Pisum sativum (clone HW39-4) gene fragment	0.05
		, , , , , , , , , , , , , , , , , , , ,	
		Lacerta vivipara 12S ribosomal RNA gene, partial sequence; tRNA-	
i		Val gene, complete sequence; and 16S ribosomal RNA gene, partial	
3175	AF206594	sequence; mitochondrial genes for mitochondrial products	0.69
3176	XM_037847	Homo sapiens neurexophilin 3 (NXPH3), mRNA	6.7
		Rattus norvegicus T:G mismatch thymine glycosylase mRNA,	
3177	AY026945	complete cds	1.6
3178	XM_028009	Homo sapiens KIAA1563 protein (KIAA1563), mRNA	2E-71
		Carsonella ruddii RNA polymerase beta subunit (rpoB) and RNA	
3179	AF250386	polymerase beta-prime subunit (rpoC) genes, partial cds	0.65
3180	AB038968	Homo sapiens CFLAR gene, exon 5, exon 6	0.24
3182	AK023569	Homo sapiens cDNA FLJ13507 fis, clone PLACE 1005085	2E-64
3183	AF240176	Mus Mus musculus MRP13 mRNA	2
3184	U20230	Human guanyl cyclase C gene, partial cds	0.2
		Ovis aries T cell receptor gene V-region, exons 1 (3' end) and 2 (5'	
3185	L18959	end)	0.64
3186	U36785	Rattus norvegicus putative pheromone receptor VN1 mRNA, complete	
3187	AF270614	cds Homo sapiens clone 5qtel_c93t7 sequence	0.73
3188	NM 004114	Homo sapiens fibroblast growth factor 13 (FGF13), mRNA	0.075
3189	AJ403239	M.musculus DNA for vimentin-binding fragment VimE7	6,7 0,076
3109	A3403233	Mus musculus, clone MGC:6551 IMAGE:2655861, mRNA, complete	0.076
3190	BC010716	cds	4.9
		Staphylococcus epidermidis strain SR1 clone step 1051e05 genomic	
3191	AF270162	sequence	0.56
		Homo sapiens hypothetical gene supported by AL137734	
3192	XM_046992	(LOC92742), mRNA	1.5
3193	AE003909	Yulella factidioca 0a5a caction 55 of 220 of the	6.2
3193	AF240176	Xylella fastidiosa 9a5c, section 55 of 229 of the complete genome  Mus Mus musculus MRP13 mRNA	6.3 1.5
3194	211 240170	Romanomermis culicivorax mitochondrial NADH dehydrogenase	1.3
		subunits 6 and 3 (ND6, ND3) cytochrome P450-like protein genes,	
3195	L08174	complete CDS	0.5
3173	200174	compact CDB	0.5

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID	-		
NO	ACCESSN	DESCRIP	P VALUE
		Mus musculus adult male corpus striatum cDNA, RIKEN full-length	
3196	AK021098	enriched library, clone:C030021G16, full insert sequence	6.5
3197	AT211627	Eimeria separata ORF470 gene, partial cds; apicoplast gene for	
3197	AF311637	apicoplast product	0.079
3198	D84549	Candida tropicalis DNA for carnitine acetyltransferase, complete cds	0.057
3199	XM_045864	Homo sapiens KIAA1804 protein (KIAA1804), mRNA	0
3200	XM_010334	Homo sapiens transducin (beta)-like 1 (TBL1), mRNA	1.9
3201	AL133154	S.pombe chromosome I cosmid c1751	0.15
			i —
3202	S75898	aromatase [Poephila guttata=zebra finches, ovary, mRNA, 3188 nt]	0.49
		Homo sapiens dystrophin (DMD) gene, deletion junction 8 between	
3203	AF047669	introns 7 and 20	0.001
			ĺ
3204	AB042652	Bos taurus mRNA for zona pellucida glycoprotein ZPB, complete cds	0.58
3205	AF147082	Homo sapiens gamma-glutamyl hydrolase gene, exons 3 through 7	0.028
3206	AE006252	Lactococcus lactis subsp. lactis IL1403 section 14 of 218 of the complete genome	
3207	Z99773	Caenorhabditis elegans cosmid H06A10, complete sequence	0.07
3207	NC 001328	Caenorhabditis elegans mitochondrion, complete genome	0.7
3208	NC_001328	Caenornaoulus elegans, muochondrion, complete genome	0.025
3209	AY037251	Arabidopsis thaliana AT4g30750/T10C21 100 mRNA, complete cds	5.9
3210	Z12834	L.esculentum gene for polyphenol oxidase	0.75
		Human DNA sequence from clone RP11-533B2 on chromosome 10,	
3211	AL391057	complete sequence [Homo sapiens]	0.07
		Helianthus annuus Dc3 promoter-binding factor-2 (DPBF-2) mRNA,	
3212	AF001454	complete cds	0.21
		Streptococcus pneumoniae transposon Tn5252 Orf28, Orf27, Orf26,	
3213	AF295925	Orf25, Orf24, Orf23, Orf22, Orf21, and Orf20 genes, complete cds	0.019
3214	AC024821	Caenorhabditis elegans cosmid Y55B1AR, complete sequence	1.7
		Homo sapiens 12q BAC RP11-946G22 (Roswell Park Cancer Institute	
3215	AC018655	Human BAC Library) complete sequence	0.0000001
		Homo sapiens cadherin 17, LI cadherin (liver-intestine) (CDH17),	
3216	XM_043524	mRNA	2E-52
		Homo sapiens phosphoinositide-3-kinase, catalytic, beta polypeptide	
3217	XM_051566	(PIK3CB), mRNA	0.025
3218	X84742	M.musculus GPX3 gene	0.079
3219	AL445527	Human DNA sequence from clone RP11-115C2 on chromosome 1, complete sequence [Homo sapiens]	
3219	AL443327		5.4
3220	AF254792	Danio rerio Period3 circadian clock protein (Per3) mRNA, complete cds	3.8
3220	AF234/92	Chlamydophila pneumoniae AR39, section 53 of 94 of the complete	3.8
3221	AE002225	genome genome AR39, section 53 of 94 of the complete	0.12
3221	AEUUZZZ3	Benome	U.12

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ļ		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Homo sapiens similar to hypothetical protein (H. sapiens)	
3222	XM_048692	(LOC92998), mRNA	2E-15
		Clavulina cinerea 18S ribosomal RNA gene, partial sequence; internal	
		transcribed spacer 1, 5.8S ribosomal RNA gene and internal	
3223	AF335456	transcribed spacer 2, complete sequence; and 28S ribosomal RNA gene, partial sequence	0.007
3223	AI-333430	Glycine max chitinase class I (Chia1) and hypothetical protein genes,	0.037
3224	AF335589	complete cds	3.3
5221	TH SSCOO	Escherichia coli O157:H7 EDL933 genome, contig 1 of 3, section 54	3.3
3225	AE005230	of 155	5.2
		Homo sapiens MASP gene for mannose binding protein-associated	3,2
3226	AB010822	protease, exon 1	7E-13
3227	XM_052139	Homo sapiens interleukin 13 receptor, alpha 1 (IL13RA1), mRNA	0.007
	-	Toxoplasma gondii TgMIC10 precursor (MIC10) mRNA, complete	
3228	AF293654	cds	0.15
3229	AJ388514	Canis familiaris mRNA for partial Ribosomal protein S14	0.38
3230	U80814	Caenorhabditis elegans cosmid B0547, complete sequence	0.22
3231	NM_013890	Mus musculus f-box and WD-40 domain protein 2 (Fbxw2), mRNA	0.29
2222	A D040411	Homo sapiens strg gene for striatum-specific G potein-coupled	
3232 3233	AB042411	receptor, complete cds	0.000009
3233	AF151527	Hyphantria cunea serpin gene, complete cds  M. musculus (SRP9) signal recognition particle subunit mRNA,	0.001
3234	X78305	1116bp signal recognition particle subunit mRNA,	0.047
3231	21.10505	11100	0.047
3235	NM 018830	Mus musculus N-acylsphingosine amidohydrolase 2 (Asah2), mRNA	0.18
		Apocryptophagus sp. MSPAR4 cytochrome oxidase I (COI) gene,	0.10
3236	AF302060	partial cds; mitochondrial gene for mitochondrial product	0.007
		Arabidopsis thaliana genomic DNA, chromosome 5, TAC	
3237	AB024025	clone:K16F13	0.012
3238	U25810	Bos taurus lysozyme (LZ) gene, complete cds	0.004
3239	X95363	C.annuum gamma thionin gene	0.15
		Plasmodium berghei carbamoyl phosphate synthetase II gene, partial	
3240	AF286897	cds	0.15
3241	AF290920	Procambarus clarkii microsatellite PcIG-03 sequence	3.7
3242	AT00<000	Lactococcus lactis subsp. lactis IL1403 section 142 of 218 of the	
3242	AE006380 U66913	complete genome	2.2
3243	000913	Dictyostelium discoideum ORF DG1040 gene, partial cds Mouse genes for transfer RNA-Ala, tRNA-Ile, tRNA-Pro and tRNA-	0.22
3244	Y00460	Mouse genes for transfer RNA-Ata, tRNA-Ite, tRNA-Pro and tRNA- Lys	0.026
	100400	Fruitfly myosin alkali light chain (Mlc1) gene	0.020
		`supercontractile/tubular muscle and indirect flight muscle isoforms';	
3245	L08052	complete cds's	0.67
		Dictyostelium discoideum ubiquitin-conjugating enzyme protein UbcC	0.01
3246	AF076597	(ubcC) mRNA, complete cds	0.58
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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
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NO	ACCESSN	DESCRIP	P VALUE
3247	U67518	Methanococcus jannaschii section 60 of 150 of the complete genome	0.003
	007010	Dictyostelium discoideum histidine kinase DhkH (dhkH) gene, partial	0,003
3248	AF362370	cds	6
		Fugu rubripes sex comb on midleg-like 2 protein (SCML2) gene,	
3249	AF146688	complete cds	0.026
		Mus musculus adult male tongue cDNA, RIKEN full-length enriched	
3250	AK009952	library, clone:2310057C01, full insert sequence	0.07
3251	AB005803	Homo sapiens DNA for histidine-rich glycoprotein, complete cds	0.078
3252	AY021484	Oryza sativa microsatellite MRG3809 containing (TA)X27, closest to marker G227, genomic sequence	
3232	A1021404	Homo sapiens DNA from chromosome 19-cosmid f19399 (~17 kb	2.2
3253	AD000833	EcoRI restriction fragment)	0.024
3254	AB048982	Macaca fascicularis brain cDNA, clone:QnpA-13067	0.38
3255	AL023839	Caenorhabditis elegans cosmid Y39A1C, complete sequence	0.026
3256	NC 002052	Tomato spotted wilt virus RNA-L, complete sequence	0.22
3257	AK022137	Homo sapiens cDNA FLJ12075 fis, clone HEMBB1002425	0.64
3258	U44047	Xenopus laevis MCM2 mRNA, complete cds	0.24
		H.sapiens CpG island DNA genomic Mse1 fragment, clone 51d3,	
3259	Z61385	forward read cpg51d3.ft1a	5E-21
		Homo sapiens dentin sialophosphoprotein precursor (DSPP) gene,	
3260	AF163151	complete cds	0.003
		Montinia caryophyllacea ribulose 1,5-bisphosphate carboxylase large	
3261	L11194	subunit (rbcL) gene, partial cds; chloroplast gene for chloroplast	
3201	L11194	Homo sapiens, clone MGC:11357 IMAGE:3954507, mRNA,	6.4
3262	BC002928	complete cds	3E-46
3263	XM_002837	Homo sapiens acid phosphatase, prostate (ACPP), mRNA	2.3
5205	1211_002037	Hordcum vulgare putative c-myb-like transcription factor (MYB3R-1)	2.3
3264	AF189787	gene, partial cds	0.61
3266	AB048353	Paramecium caudatum mRNA for hsp60, partial cds	1.8
		Arabidopsis thaliana acyl-CoA binding protein (ACBP) gene,	
3267	U75274	complete cds	0.69
3268	AF034077	Equus caballus alpha-1-antitrypsin (Spi2) gene, complete cds	0.077
		Human DNA sequence from clone CTA-191D12 on chromosome	
		22q13.1 Contains two exons of the APOL2 gene for apolipoprotein L	
3269	AL031426	2, ESTs and GSSs, complete sequence [Homo sapiens]	0.000001
3270	AJ007958	Thiobacillus sp. plasmid pT3.2I including adh and repA genes, partial	0.001
		Mus musculus RIKEN cDNA 2700016E08 gene (2700016E08Rik),	
3271	NM_025978	mRNA	1E-17
	*****	Homo sapiens similar to gene with multiple splice variants near HD	
3272	XM_018403	locus on 4p16.3 (H. sapiens) (LOC91016), mRNA	5.5
3273	AJ238613	Gallus gallus mRNA for teneurin-1	0.57
3274	XM_033841	Homo sapiens FGFR1 oncogene partner (FOP), mRNA	0.55

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			Т
NO	ACCESSN	DESCRIP	P VALUI
3275	NM_014601		0.065
3276	AK021659	Homo sapiens cDNA FLJ11597 fis, clone HEMBA1003856	9E-13
3277	AB054644	Sarcophaga peregrina gene for lectin, 5' upstream region and partial cds	0.045
3278	AK010362	Mus musculus ES cells cDNA, RIKEN full-length enriched library, clone:2410003J06, full insert sequence	5.5
3279	AF282974	Plasmodium falciparum cysteine protease falcipain-3 gene, complete cds	1.8
3280	U60135	Arabidopsis thaliana serine/threonine protein phosphatase 2A-3 catalytic subunit gene, complete cds	0.024
3281	AF365086	Martiodendron parviflorum tRNA-Leu (trnL) gene, partial intron sequence; chloroplast gene for chloroplast product	6.5
3282	AE002140	Ureaplasma urealyticum section 41 of 59 of the complete genome	0.22
3283	Z49124	S.oleracca chloroplast secA mRNA	0.074
3284	AE002353	Chlamydia muridarum, section 80 of 85 of the complete genome	4.4
3285	XM_011989	Homo sapiens ELKL motif kinase (EMK1), mRNA	7E-54
3286	AJ252287	Plasmodium falciparum partial msp3 gene for merozoite surface protein 3	0.68
3287	AK018224	Mus musculus adult male medulla oblongata cDNA, RIKEN full- length enriched library, clone:6330531101, full insert sequence	2
3288	X91123	Homo sapiens partial SLC6A2 gene for norepinephrine transporter, exon 5	7.5
3289	AB045007	Mus musculus GNB2 gene for guanine nucleotide binding protein beta2 subunit, complete cds	0.67
3290	AY017272	Peromyscus difficilis isolate TK32541 mys-9 retrotransposon and LTR, 5' sequence	5,4
3291	AL590902	S.pombe chromosome I cosmid pB8E5	0.077
3292		Homo sapiens surfactant protein A binding protein (SPAR), mRNA	0.38
3293	AK027388	Homo sapiens cDNA FLJ14482 fis, clone MAMMA1002362	0.007
2204	A Do sacos	Homo sapiens BHLHB2 gene for bHLH transcriptional factor DEC1,	
3294	AB043885	complete cds	0,068
3295	XM_028213	Homo sapiens up-regulated by BCG-CWS (LOC64116), mRNA	0.073
3296	XM_039789	Homo sapiens solute carrier family 2 (facilitated glucose transporter), member 2 (SLC2A2), mRNA	9.2
3297	AK021841	Homo sapiens cDNA FLJ11779 fis, clone HEMBA1005921	0.65
3298	M11815	Plasmid pGKL1 from killer yeast (K.lactis), complete	0.008
3299	AF166527	Zea mays flowering-time protein isoforms alpha and beta (ZmLD) gene, alternatively spliced products, complete cds	0.074
3300	XM_040556	Homo sapiens similar to hypothetical protein FLJ20127 (H. sapiens) (LOC91775), mRNA	0.22
3301	AB001569	Carrot DNA for transposon Tdc1	0.17
3302	AF326737	Mus musculus sclerostin gene, complete cds	4.7
3303	AF272001	Ebola virus subtype Zaire strain Mayinga complete genome	1.9

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#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE Homo sapiens similar to hypothetical protein FLJ22294 (H. sapiens) XM 034863 (LOC90918), mRNA 3304 0.2 3305 AC006654 Caenorhabditis elegans cosmid H09I01, complete sequence 0.023 Mus musculus adult male stomach cDNA, RIKEN full-length 3306 AK008812 enriched library, clone:2210403E17, full insert sequence 6.1 Lucilia eximia tRNA-lle gene, partial sequence: D-loop, complete sequence; and 12S ribosomal RNA, partial sequence; mitochondrial 3307 AF151387 genes for mitochondrial products 0.077 immunoglobulin epsilon chain constant region=secreted form {3' 3308 S53497 region} [human, B cell myeloma U-266, Genomic, 3198 nt] 5.5 3309 Homo sapiens stromal antigen 2 (STAG2), mRNA XM 010126 e-115 Homo sapiens MAT gene for mitochondrial acetoacetyl-CoA thiolase, 3310 D10500 1.9 Paraflabellula reniformis small subunit ribosomal RNA gene. 3311 AF293900 complete sequence 0.092 3312 NC_001530 Human papillomavirus type 47, complete genome 0.67 H. sapiens Ig lambda light chain variable region gene (34-34SWIIF32) 3313 785396 rearranged; Ig-Light-Lambda; VLambda 0.71 3314 NC '001566 Apis mellifera ligustica mitochondrion, complete genome 0.015 Plasmodium voclii blood-stage membrane protein Ag-1 mRNA. 3315 AF103869 complete cds 1.5 3316 XM 007957 Homo sapiens hypothetical protein (LOC57019), mRNA 3E-27 Homo sapiens chromosome 18 open reading frame 1 (C18orf1). 3317 XM 012723 mRNA 27 XM 046956 Homo sapiens Kallmann syndrome 1 sequence (KAL1), mRNA 3318 4.5 3319 Helicobacter pylori 26695 section 92 of 134 of the complete genome 0.068 3320 NM 017932 Homo sapiens hypothetical protein FLJ20700 (FLJ20700), mRNA 1.7 Mus musculus 8 days embryo cDNA, RIKEN full-length enriched 3321 AK017541 library, clone:5730410F24, full insert sequence 4.8 Mus musculus adult male testis cDNA, RIKEN full-length enriched 3322 AK016519 library, clone:4932411G06, full insert sequence 1.9 Trypanosoma cruzi kinetoplast minicircle ATPase subunit 6 mRNA. 3323 U38184 complete cds 0.0003 Z54181 3324 M.catarrhalis bla gene for BRO-2 0.069 Oryza sativa 3-hydroxy-3-methylglutaryl-CoA reductase gene, 3325 U43961 promoter region and complete cds 0.17 3326 AK025166 Homo sapiens cDNA: FLJ21513 fis, clone COL05778 0.23 3327 AF158727 Homo sapiens truncated properdin (PFC) gene, exons 1 and 2 0.73 Botrytis cincrea strain T4 cDNA library under conditions of nitrogen 3328 AT.116536 deprivation 0.57 3329 NC 001712 Locusta migratoria mitochondrion, complete genome 0.061

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		W. 11 At 17	
		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	77767-11	
NU	ACCESSN	DESCRIP	P VALUE
		Stanbulgesease anidemidical and a CTC but I at	
		Staphylococcus epidermidis plasmid pST6 beta-lactamase repressor BlaI (blaI) gene, partial cds; resolvase/integrase BinR (binR),	
		recombinase Sin (sin), repressor QacR (qacR), and transport protein	l
3330	AY028779	QacB (qacB) genes, complete cds; and unknown gene	
3331	AF346992	Homo sapiens mitochondrion, complete genome	0.23 1.7
5551	111310332	Traine superior intectionation, complete genome	1./
3332	NM_031389	Mus musculus ribonuclease/angiogenin inhibitor 2 (Rnh2), mRNA	1.9
		g-Benn (material), mit ur	
3333	AC079918	Homo sapiens BAC clone RP11-114K13 from 7, complete sequence	0.0002
3335	AF298207	Dictyostelium discoideum transposon thug-S, complete sequence	0.15
		Saccharomyces cerevisiae mitochondrion 21S ribosomal RNA (21S	
		rRNA) gene, 21S R2 ribosomal RNA (21S R2 rRNA) gene, and ORF	
3336∙	L36903	RI	0.024
3337	AF239663	Prunus persica ACC synthase gene, partial cds	0.066
		H.sapiens D2-1-K2G7np gene for immunoglobulin kappa chain	
3338	AJ223690	variable region	0.026
3339	AC084157	Caenorhabditis elegans cosmid Y46E12BR, complete sequence	4.4
3340	M58606	B.subtilis threonine deaminase (ILVA) gene, complete cds	0.62
3341	NC_001648	Cassava vein mosaic virus, complete genome	0.008
- 1		Caenorhabditis elegans Bristol N2 synaptic vesicle acetylcholine	
3342	U09277	transporter (unc-17) gene, complete cds	0.024
3343	U52821	Mus musculus prion (PrP) gene, exons 1 and 2	1.9
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
3344	AK014936	library, clone:4921521D15, full insert sequence	2
3345	AE007447	g	_
3343	AE007447	Streptococcus pneumoniae section 130 of 194 of the complete genome Mus musculus adult male testis cDNA, RIKEN full-length enriched	5
3346	AK005820	library, clone:1700010F03, full insert sequence	0.10
3347	AB011681	Danio rerio mRNA for ZOR-2, complete cds	1.9
3341	ADOTTOST	Mus musculus Chetk-alpha gene for choline/ethanolamine kinase-	1.9
3348	AB030616	alpha, exon 1 and 5'-flanking region	2E-10
3340	110000010	Staphylococcus epidermidis strain SR1 clone step.1002e12 genomic	26-10
3349	AF269359	sequence	0.53
			0.55
3350	AF063004	Trypanosoma rotatorium trans-spliced leader gene, partial sequence	1.9
		Plasmodium falciparum chromosome 2, section 48 of 73 of the	
3351	AE001411	complete sequence	0.008
3352	AF240496	Mus musculus clone 148d4 Naip5 gene, partial sequence	0.56
3353	XM_004722	Homo sapiens hypothetical protein FLB6421 (FLB6421), mRNA	2.2
1		Mycoplasma fermentans DNA polymerase III subunit (dnaH), YabC	
		protein (yabC), orf550, lipoprotein MALP-404 precursor (malP),	
- 1		insertion sequence IS1630 transposase (tnpA) genes, complete cds;	
3354	AF100324	P78 ATP-binding cassette transport operon, complete sequence;>	2.1
3355	NM_019684	Mus musculus serine/threonine kinase 23 (Stk23), mRNA	0.23

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WO0214500 [fle //E /WO0214500 opc]

WO 02/14500

#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE 3356 AB004856 Buchnera aphidicola mRNA for homoserine kinase, partial cds 1.9 3357 AF328428 Homo sapiens clone 1ptel_2109Bf&Br sequence 2E-11 Homo sapiens hypothetical gene supported by AF308287; BC009516 3358 XM 041166 (LOC91860), mRNA 0.024 Lactococcus lactis subsp. lactis IL1403 section 191 of 218 of the 3359 AE006429 complete genome 0.24 HIV-1 clone 8 isolate BV from Milan, envelope glycoprotein, V1-V5 3360 U95455 region (env) gene, partial cds 0.026 Homo sapiens PEF protein with a long N-terminal hydrophobic 3361 XM 001775 domain (peffin) (PEF), mRNA 0.45 3362 AF100670 Caenorhabditis elegans cosmid M4 3363 AB036737 Homo sapiens mRNA for RERE, complete cds 3364 AE001030 Archaeoglobus fulgidus section 77 of 172 of the complete genome 1.5 Homo sapiens hypothetical gene supported by AK027658 3365 XM 046602 (LOC92679), mRNA 2 3366 XM 005889 Homo sapiens kinesin-like 1 (KNSL1), mRNA 0.21 Lycopersicon esculentum MADS-box transcription factor jointless 3367 AF275345 gene, complete cds 1.2 NC 000857 | Ceratitis capitata complete mitochondrial genome 3368 0.022 Human PCR Fragment of Cosmid g1346a274 from 7q31.3, complete 3369 AC002048 sequence [Homo sapiens] 1E-16 3371 AB016611 Enterobacter cloacae ampC gene, complete cds 0.62 Mus musculus RIKEN cDNA 2810036K01 gene (2810036K01Rik). 3372 NM_026040 e-138 Human immunodeficiency virus type 1, partial proviral POL gene, rt 3373 AJ252860 region, isolate CL 0.25 3374 XM_043434 Homo sapiens villin-like (VILL), mRNA 1.8 Drosophila melanogaster genomic scaffold 142000013385952. 3375 AE002776 complete sequence 0.19 Homo sapiens tumor necrosis factor receptor superfamily, member 8 XM 030198 3376 (TNFRSF8), mRNA 0.76 3377 X65450 C.canis mRNA for chloride channel 6.3 Homo sapiens hypothetical protein DKFZp434F1719 3378 NM 032248 (DKFZp434F1719), mRNA 1E-47 Carnobacterium piscicola transposase, bacteriocin, histidine protein kinase, ATP dependent transloactor, accessory protein, and 3379 L47121 carnobacteriocin B2 genes, complete eds 0.18 3380 L26287 Schistosoma mansoni SMDR2 gene, complete cds 0.66 3381 AF125130 Nemadactylus macropterus clone 15,1 microsatellite sequence 0.072 Oryzias latines olgc4 gene for membrane guanylyl cyclase OIGC4, 5' 3383 AB040099 flanking sequence and partial cds 0.22 AF375398 3384 Arabidopsis thaliana At1g51660/F19C24 26 gene, complete cds 6 HIV-1 clone TV004G24 from South Africa gag protein (gag) gene, 3385 AF391256 complete cds 0.23 3386 X91865 A.thaliana asp5 gene 2 1

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	-	Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID	ł		
NO	ACCESSN	DESCRIP	P VALU
		Homo sapiens variable charge protein on X with eight repeats (VCX-	
3410	XM_047898		5.9
	*D * 0 ***	Homo sapiens TATA box binding protein (TBP)-associated factor,	
3411	XM_047003		0,7
3412	D86566	Human DNA for NOTCH4, partial cds	0.025
3413	AE260201	Arabidopsis thaliana unknown protein (T8B10_70/AT3g60410)	
3414	AF360301 AJ404228	mRNA, complete cds Saccharomyces douglasii mitochondrial tRNA gene cluster	0.64
3414	AJ404228	Sacciaroniyees douglash mitochondriai tRNA gene cluster	0.002
3415	XM 037369	Home conions hypothetical matrix TX 100000 (TX 100000) TX 1	
3413	ZIVI_037309	Homo sapiens hypothetical protein FLJ20693 (FLJ20693), mRNA Escherichia coli O157:H7 EDL933 genome, contig 1 of 3, section 54	9E-14
3416	AE005230	of 155	
3710	AL003230	O1 133	6,2
. !		Thomas DNA services for the BDs seedle	1
		Human DNA sequence from clone RP5-839B11 on chromosome 20. Contains the first coding exon of the gene for a novel protein with a	ļ
		Kunitz/Bovine pancreatic trypsin inhibitor domain and WAP-type	
3417	AL121778	(Whey Acidic Protein) 'four-disulfide core' domains and an e>	0.08
		Oryza sativa microsatellite MRG3085 containing (TA)X13, closest to	0.08
3418	AY020760	marker R2976, genomic sequence	6,8
		Homo sapiens mRNA; cDNA DKFZp434L098 (from clone	0.8
3419	AL122122	DKFZp434L098)	1.8
		Xenopus laevis early growth response protein 1 (egr1) gene, partial	-110
3420	AF250346	cds	0,077
3421	XM_044852	Homo sapiens KIAA1572 protein (KIAA1572), mRNA	0.7
3422	XM_049057	Homo sapiens G protein-coupled receptor 49 (GPR49), mRNA	7
		H.sapiens D2-1-K2G7np gene for immunoglobulin kappa chain	
3423	AJ223690	variable region	0.028
		Anas platyrhynchos mIgM gene (partial), mIgA gene (partial) and	
3424	AJ314752	IgA gene (partial), clone 13.1	0.072
3425	U51739		
	031737	Ipomoea trifida secreted glycoprotein 2 (ISG2) mRNA, complete cds	0.64
3426	S66752	coagulation factor IX {3' region} [human, Genomic Mutant, 3130 nt]	0.026
3426 3427	S66752 AE001122	coagulation factor IX {3' region} [human, Genomic Mutant, 3130 nt] Borrelia burgdorferi (section 8 of 70) of the complete genome	0.026
3426	S66752	coagulation factor IX {3' region} [human, Genomic Mutant, 3130 nt]	0.026
3426 3427 3428	\$66752 AE001122 AJ233850	coagulation factor IX (3¹ region). [human, Genomic Mutant, 3   30 m] Borrelia burgdorferi (section 8 of 70) of the complete genome Raitus norvegicus microsatellite sequence clone 27E9	0.026 0.72 2.1
3426 3427 3428 3429	S66752 AE001122 AJ233850 L35664	coagulation factor IX (3' region) [human, Genomic Mutant, 3130 m] Borrelia burgdorferi (section 8 of 70) of the complete genome Rattus norvegicus microsatellite sequence clone 27E9  Homo sapiens (subclone H8 8 15 from P1 35 H5 C8) DNA sequence	0.026 0.72 2.1 0.079
3426 3427 3428	S66752 AE001122 AJ233850 L35664	coagulation factor IX {3' region} [human, Genomic Mutant, 3130 nt] Borrelia burgdorferi (section 8 of 70) of the complete genome Rattus norvegicus microsatellite sequence clone 27E9  Homo sapiens (subclone H8 8 f5 from P1 35 H5 C8) DNA sequence Callithrix geoffroyi epsiton-globin gene, partial cds	0.026 0.72 2.1
3426 3427 3428 3429	S66752 AE001122 AJ233850 L35664	coagulation factor IX [3' region] [human, Genomic Mutant, 3130 m] Borrelia burgdorferi (section 8 of 70) of the complete genome Rattus norvegicus microsatellite sequence clone 27E9 Homo sapiens (subclone H8 8 f5 from Pl 35 H5 C8) DNA sequence Callithris geoffryi epsilon-globin gene, partial cds Human DNA sequence from clone LL22N(O1-86D10 on chromosome	0.026 0.72 2.1 0.079
3426 3427 3428 3429 3430	S66752 AE001122 AJ233850 L35664 U97029	coagulation factor IX (3' region) [human, Genomic Mutant, 3130 m] Borrelia burgdorferi (section 8 of 70) of the complete genome Rattus norvegicus microsatellite sequence clone 27E9  Homo sapiens (subclone H8 8, E5 from P1 35 H5 C8) DNA sequence Callithris geoffroyi epsilon-globin gene, partial cds Human DNA sequence from clone LL22NC01-86D10 on chromosome 22 Contains part of the SYN3 gene for synapsin III and ESTs,	0.026 0.72 2.1 0.079 2.1
3426 3427 3428 3429	\$66752 AE001122 AJ233850 L35664 U97029	coagulation factor IX (3' region) [human, Genomic Mutant, 3130 m] Borrelia burgdorferi (section 8 of 70) of the complete genome Rattus norvegicus microsatellite sequence clone 27E9 Homo sapiens (subclone H8 8 J5 from P1 35 H5 C8) DNA sequence Callithrix geoffroyi epsilon-globin gene, partial cds Human DNA sequence from clone LL22NC01-8cD10 on chromosome 22 Contains part of the SYN3 gene for synapsin III and ESTs, complete sequence [Homo sapience [Homo sapience]]	0.026 0.72 2.1 0.079
3426 3427 3428 3429 3430 3431	\$66752 AE001122 AJ233850 L35664 U97029 Z82181	coagulation factor IX [3' region] [human, Genomic Mutant, 3130 m] Borrelia burgdorfert (section 8 of 70) of the complete genome Rattus norvegicus microsatellite sequence clone 27E9 Homo sapiens (subclone H8 8, f5 from P1 35 H5 C8) DNA sequence Callithrix geoffroyi epsilon-globin gene, partial cds Human DNA sequence from clone L122NC01-8E010 on chromosome 22 Contains part of the SYN3 gene for synapsin III and ESTs, complete sequence [Homo sapiens] Macaca fascicularis brain cDNA clone-QtrA-10780, full insert	0.026 0.72 2.1 0.079 2.1
3426 3427 3428 3429 3430 3431	S66752 AE001122 AJ233850 L35664 U97029 Z82181 AB066544	coagulation factor IX (3' region) [human, Genomic Mutant, 3130 m] Borrelia burgdorferi (section 8 of 70) of the complete genome Rattus norvegicus microsatellite sequence clone 27E9  Homo sapiens (subclone H8 8 15 from P1 35 H5 C8) DNA sequence Callithrix geoffroyi epsilon-globin gene, partial cds Human DNA sequence from clone LL22NC01-8cD10 on chromosome 22 Contains part of the SYNS gene for synapsin III and ESTs, complete sequence [Homo sapiens] Macaca fascicularis brain cDNA clone-QtrA-10780, full insert sequence	0.026 0.72 2.1 0.079 2.1 0.71
3426 3427 3428 3429 3430 3431	S66752 AE001122 AJ233850 L35664 U97029 Z82181 AB066544	coagulation factor IX [3' region] [human, Genomic Mutant, 3130 m] Borrelia burgdorfert (section 8 of 70) of the complete genome Rattus norvegicus microsatellite sequence clone 27E9 Homo sapiens (subclone H8 8, f5 from P1 35 H5 C8) DNA sequence Callithrix geoffroyi epsilon-globin gene, partial cds Human DNA sequence from clone L122NC01-8E010 on chromosome 22 Contains part of the SYN3 gene for synapsin III and ESTs, complete sequence [Homo sapiens] Macaca fascicularis brain cDNA clone-QtrA-10780, full insert	0.026 0.72 2.1 0.079 2.1

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#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE Kladothrips rugosus cytochrome oxidase subunit I (COI) gene. 3435 1193519 mitochondrial gene encoding mitochondrial protein, partial cds 0.77 3437 1123451 Caenorhabditis elegans cosmid ZK75 0.48 3438 BC003415 Homo sapiens, clone IMAGE:3451161, mRNA 0.41 3439 AK024526 Homo sapiens cDNA: FLJ20873 fis, clone ADKA02669 3.3 Neisseria meningitidis serogroup B strain MC58 section 190 of 206 of 3440 AE002548 the complete genome 06 Anopheles minimus strain PT1 cytochrome oxidase subunit 1 (COI) 3441 AF116832 gene, partial cds; mitochondrial gene for mitochondrial product 1.5 3442 AB055323 Macaca fascicularis brain cDNA, clone:QflA-10308 16 Mus musculus 0 day neonate head cDNA, RIKEN full-length enriched AK014667 3443 library, clone:4833408P15, full insert sequence 7E-25 3444 U80753 Homo sapiens CAGL79 mRNA, partial cds 0.048 AJ275304 Cicer arietinum mRNA for putative ABA-responsive protein 3445 1.4 3446 AF336131 Lumpy skin disease virus strain Neethling, 3' partial sequence 0.3 HIV-1 isolate LBLA010 from Liberty City, Florida, envelope 3447 U11476 glycoprotein (env) gene, partial cds 16 Homo sapiens hypothetical gene supported by AL137476 3448 XM 045582 (LOC92530), mRNA 6E-13 3449 AF030199 Mus musculus type 1 sigma receptor gene, complete cds 0.63 Myzopoda aurita 12S ribosomal RNA gene, complete sequence: tRNA-Val gene, complete sequence; and 16S ribosomal RNA gene, complete 3450 AF345926 sequence; mitochondrial genes for mitochondrial products 0.13 3451 NM 008715 Mus musculus Notch2-like (Notch21), mRNA 0.17 3452 AF067794 Nicotiana tabacum meristem-specific promoter sequence 0.1 AJ287175 3453 Mucor indicus partial act-1 gene for actin 3454 U89439 Bos taurus ubiquitin-like protein mRNA, complete cds 0.000003 Xenopus laevis early growth response protein 1 (egr1) gene, partial 3455 AF250346 0.072 3456 Z73587 S.cerevisiae chromosome XVI reading frame ORF YPL231w 3457 Macaca fascicularis brain cDNA, clone:OflA-14233 AB055310 3458 Y16090 Daucus carota Susy*Dc1 gene 0.13 AE003868 3459 Xylella fastidiosa 9a5c, section 14 of 229 of the complete genome 0.16 3460 AF281912 Homo sapiens clone 16atel c89bt7 sequence 5E-36 3461 AF126541 Ateline herpesvirus 3 H-DNA terminal repeat unit, complete sequence 0.023 Mus musculus 10 days embryo cDNA. RIKEN full-length enriched 3462 AK011178 library, clone:2600010E01, full insert sequence 1.5 Arabidopsis thaliana unknown protein (F1B16.11) mRNA, complete 3463 AF370226 3464 X94768 H.sapiens RP3 gene (XLRP gene 3) 3465 AB029068 Mus musculus gene for Ser/Thr kinase KKIAMRE, exon 1, 2, 3, 4, 5 0.57

3485

3486

3487

3488

3489

XM 032347

AF120475

U81676

AL596023

U67210

0

0.0008

0.31

0.016

0.056

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#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEQ ID NO ACCESSN DESCRIP PVALUE 3466 X84742 M.musculus GPX3 gene 0.038 Homo sapiens, Similar to RIKEN cDNA 4931428D14 gene, clone 3467 BC010011 MGC:15407 IMAGE:4309613, mRNA, complete cds 0.22 Legionella longbeachae spectinomycin 3' adenylyltransferase (aadA). possible transcriptional regulatory protein (IrpR), and possible sensor 3468 AF288536 kinase protein (lskS) genes, complete cds; and unknown genes 1 Legionella longbeachae spectinomycin 3' adenylyltransferase (aadA), possible transcriptional regulatory protein (IrpR), and possible sensor 3469 AF288536 kinase protein (lskS) genes, complete cds; and unknown genes 1.8 Papilio machaon mitochondrial ND5 gene for NADH dehydrogenase AB013150 3470 subunit 5, partial cds 5.9 3471 XM 028011 Homo sapiens CGI-145 protein (LOC51028), mRNA 0.63 3472 XM 052069 Homo sapiens actin related protein (MGC15664), mRNA 5.7 Human DNA sequence from clone RP11-533E16 on chromosome 6. 3473 AL449163 complete sequence [Homo sapiens] 0.00001 3474 X96685 B.burgdorferi cell division genes 1.9 3475 AF179589 Rana tigrina rugulosa glucagon receptor (GluR) mRNA, complete cds 0.19 Homo sapiens partial PIK3CB gene for phosphatidylinositol 3-kinase 3476 AJ297549 catalytic subunit p110beta, exons 1-2 and joined CDS 0.87 3477 XM 032707 Homo sapiens KIAA0630 protein (KIAA0630), mRNA 3478 U25032 Caenorhabditis elegans par-3 mRNA, complete eds 3479 U89283 Biomphalaria glabrata myoglobin gene, complete cds 3.2 Strongylocentrotus purpuratus ADAM precursor, mRNA, complete 3480 AY029303 5.6 Mus musculus V186.2 gene for immunoglobulin heavy chain, 3481 AB043223 clone:L1G6001 7.5 Danio rerio transmembrane receptor Roundabout1 (robo1) mRNA. 3482 AF304130 complete cds 1.8 3483 AF386077 Homo sapiens interleukin 17B (IL17B) gene, complete cds 0.65 3484 U39889 Bos taurus Y-chromosome specific genomic sequence Homo sapiens region containing TLS-associated serine-arginine protein 1; TLS-associated serine-arginine protein 2; TLS-associated

TLS-associated serine-arginine protein 1: TLS-associa>

complete cds, alternatively spliced

complete sequence [Homo sapiens]

Human clone HS2.16 Alu-Ya5 sequence

partial sequence

scrine-arginine protein 2; TLS-associated serine-arginine protein 2;

Unidentified cubacterium clone vadinBC27 16S ribosomal RNA gene.

Human DNA sequence from clone RP11-142J18 on chromosome X,

Mus musculus myelin-associated oligodendrocytic basic proteins MOBP170 and MOBP69 (Mobp) gene, exons 3 through 5 and

3514

3515

AE002304

AF083830

0.2

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#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID ACCESSN NO DESCRIP P VALUE Homo sapiens similar to ret finger protein 2; candidate tumor 3490 XM 007196 suppressor involved in B-CLL (H. sapiens) (LOC93519), mRNA 0.0000008 Human DNA sequence from clone SC22CB-23F1 on chromosome 3491 22q13.31-13.33 Contains a GSS, complete sequence [Homo sapiens] AL049568 0.074 3492 X94755 A,thaliana mRNA for SNF1-related ser/thr protein kinase (1852 bp) 0.15 Human DNA sequence from clone RP13-459H7 on chromosome 10, 3493 AL392109 complete sequence [Homo sapiens] 1.8 immunoglobulin epsilon chain constant region-secreted form {3' 3494 S53497 region) [human, B cell myeloma U-266, Genomic, 3198 nt] 3.2 3495 X16027 R norvegicus gene encoding alkaline phosphatase, exon 2 0.0009 Buchnera aphidicola plasmid pTrpEG isolate Mor26ps anthranilate 3496 AF213918 synthase component I (trpE) pseudogene, partial sequence 0.004 Streptococcus pyogenes M1 GAS strain SF370, section 20 of 167 of 3497 AE006491 the complete genome 0.33 3498 Z73587 S.cerevisiae chromosome XVI reading frame ORF YPL231w 0.089 3499 AK023040 Homo sepiens cDNA FLJ12978 fis. clone NT2RP2006321 2E-62 Homo sapiens integrin alpha-2 subunit (ITGA2) gene, ITGA2-1 3500 AF062039 allele, partial cds 0.65 3501 Z77974 H.sapiens flow-sorted chromosome 6 HindIII fragment, SC6pA5D12 0.036 3502 M13934 Human ribosomal protein S14 gene, complete cds 0.54 Mus musculus RIKEN cDNA 2010106G01 gene (2010106G01Rik), 3503 NM 023220 mRNA 1.1 AF140513 3504 Homo sapiens clone 254LL2 genomic sequence 0.0004 Homo sapiens mRNA; cDNA DKFZp762O1615 (from clone 3505 AL359558 DKFZp762O1615) 0.019 3506 XM 048768 Homo sapiens tetratricopeptide repeat domain 3 (TTC3), mRNA 0.00009 AF074994 Homo sapiens full length insert cDNA YH91C06 0.65 3508 NC 001338 Sulfolobus virus 1, complete genome 0.008 3509 U67502 Methanococcus jannaschii section 44 of 150 of the complete genome 0.21 Schizosaccharomyces pombe Byr1-binding protein Bob1 (bob1) gene. AY035821 complete cds 0.19 3510 3511 AF275842 Plasmodium falciparum clone b.2 PfEMP1 (var) mRNA, partial cds 2 AJ294714 Pichia anomala leu2 gene for beta-isopropylmalate dehydrogenase 0.19 3512 Human DNA sequence from clone RP3-397P18 on chromosome 6, 3513 AL391240 complete sequence [Homo sapiens] 3.4

(CACNA2D1) gene, exon 14

Chlamydia muridarum, section 35 of 85 of the complete genome Homo sapiens dihydropyridine receptor alpha 2 subunit

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	,
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
110	riccabbit	Escherichia coli O157:H7 EDL933 genome, contig 1 of 3, section 46	PVALUE
3516	AE005222	of 155	1.7
0515	710 4000	Escherichia coli 9142-88 cytolethal distending toxin (cdtA, cdtB, and	
3517	U04208	cdtC) genes, complete cds	0.008
3518	Y07738	M.musculus gene for vimentin	0.65
3519	AF304091	Sotalia fluviatilis alpla lactalbumin gene, partial cds	2
3520	AF270356	Staphylococcus epidermidis strain SR1 clone step.4025c09 genomic sequence	0.52
		Bombus pascuorum cytochrome b (cytb) gene, mitochondrial gene	ł
3521	AF017516	encoding mitochondrial protein, partial cds	1.8
3523	AF098919	Gallus gallus alpha-globin gene domain 5' region	2
3524	XM_047517	Homo sapiens Kruppel-like factor 4 (gut) (KLF4), mRNA	1E-39
3525	NC_001626	Petromyzon marinus mitochondrion, complete genome	0.021
		Bemisia tabaci biotype Q 16S ribosomal RNA gene, partial sequence;	
3526	AF246647	mitochondrial gene for mitochondrial product	1.9
		Human DNA sequence from clone RP3-397P18 on chromosome 6,	
3527	AL391240	complete sequence [Homo sapiens]	3.1
3528	M32779	AcNPV with an S.frugiperda insertion element IFP2.2	1.9
3529	L44121	Homo sapiens (clone pHK2.1D) CMT1A gene, repeat unit	6E-44
		Arabidopsis thaliana GTP-binding protein (ARAC5) mRNA,	
3530	U52350	complete cds	0.007
3531	X80908	L.esculentum gene for fruit ripening polygalacturonase	0.65
		Candidatus Carsonella ruddii natural-host Blastopsylla occidentalis	
1 1		ATP synthase alpha subunit (atpA) gene, partial cds; ATP synthase	
		gamma subunit (atpG) gene, complete cds; and ATP synthase beta	
3532	AF267203	subunit (atpD) gene, partial cds	0.22
	AT001000	Plasmodium falciparum chromosome 2, section 16 of 73 of the	
3533	AE001379	complete sequence	0.25
2504	70 ( 00/100	Homo sapiens hypothetical protein DKFZp761J1523	1
3534	XM_034129 AE002304	(DKFZp761J1523), mRNA	6
3536	AK024082	Chlamydia muridarum, section 35 of 85 of the complete genome	1.8
3536	AK024082	Homo sapiens cDNA FLJ14020 fis, clone HEMBA 1002508	1.7
3537	U67547	Methanococcus januaschii section 89 of 150 of the complete genome	0.22
3538	AF215629	Crypteronia paniculata chloroplast rpl16 gene, intron sequence	0.024
3539	X03853	Pea chloroplast gene for ribulose 1,5-bisphosphate carboxylase (rbcl)	0.008
3540	X85055	B.taurus cosmid-derived microsatellite DNA (clone IDVGA-40)	0.0009
		Mus musculus adult female placenta cDNA, RIKEN full-length	
3541	AK005508	enriched library, clone:1600021C16, full insert sequence	0.63
3542	Y16262	Daucus carota mRNA for neutral invertase	0.21
		Human DNA sequence from clone RP11-79A21 on chromosome X,	
3543	AL513491	complete sequence [Homo sapiens]	0.56
3544	XM 036298	Homo sapiens hypothetical gene supported by AL442095 (LOC91124), mRNA	0.02
3545		Homo sapiens KIAA0118 protein (KIAA0118), mRNA	0.46
3545		Province surprise restrict to protein (ICINAOTTO), INCAN	0.40

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Oryza sativa microsatellite MRG3085 containing (TA)X13, closest to	
3546	AY020760	marker R2976, genomic sequence	5,4
		Mycoplasma pulmonis LipB (lipB) gene, complete cds; and VsaG	
3547	AF198037	(vsaG) and VsaE (vsaE) genes, partial cds	0.008
3548	X62658	E.faecalis plasmid pAD1 sca1 gene and orfy	1.7
1 1		Emericella nidulans putative zinc finger protein (flbC) gene, complete	
3549	AF083468	cds	0.18
3550	AF315801	Homo sapiens chromosome Y AHCP pseudogene, complete sequence	1.9
3551	AC007042	Homo sapiens clone RP11-399H17, complete sequence	6
		Mycoplasma pulmonis LipB (lipB) gene, complete cds; and VsaG	
3552	AF198037	(vsaG) and VsaE (vsaE) genes, partial cds	0.008
3553	AK025090	Homo sapiens cDNA: FLJ21437 fis, clone COL04285	0
(		K.lactis APA2 gene for tetraphosphatase, QCR7 gene, bc1 complex	
3554	X76027	subunit VII	0.4
3555	Z78713	H.sapiens flow-sorted chromosome 6 HindIII fragment, SC6pA15A11	0.61
	*****	Streptococcus zooepidemicus zoocin A immunity factor (zif) and	
3556	U50357	zoocin A endopeptidase (zooA) genes, complete cds	0.13
2555	TT00 00 1	Homo sapiens green cone photoreceptor pigment gene, 5' flanking	
3557	U93721	region	1.4
3558	AJ291489	Paracentrotus lividus mRNA for fibrosurfin (surfin2656 gene)	3.7
3559	3.51.0000	Plasmid Collb-p9 colicin Ib structural and immunity genes, complete	
3559	M13820	cds Human DNA sequence from cosmid L129H7, Huntington's Disease	0.2
3560	Z54147	Region, chromosome 4p16.3 contains CpG island	2
3561	AJ301616	Homo sapiens DNC gene for deoxynucleotide carrier, exons 1-9	0.17
3301	AJ301010	Human immunodeficiency virus type 1 defective gag gene, clone 3-13,	0.17
3562	U29403	truncated cds	0.63
3302	029403	Plasmodium falciparum heat shock protein hsp104 (hsp104) gene,	0.03
3563	AF090118	partial cds	0.21
3564	722707	S.scrofa spl IFN gene	0.57
3565	AF090889	Homo sapiens clone HQ0092	0.62
		Troub suprem grant - 1200	0.02
3566	XM 050705	Homo sapiens GalNAc-4-sulfotransferase 2 (GALNAC4ST-2), mRNA	0.15
-		Homo sapiens similar to 8-oxoguanine DNA glycosylase (H. sapiens)	0.15
3567	XM 016611	(LOC93577), mRNA	0,052
3568	Z35914	S.cerevisiae chromosome II reading frame ORF YBR045c	0.59
		CYP1A2=3-methylcholanthrene responsive gene [Oncorhynchus	
3569	S69278	mykiss=rainbow trout, liver, Genomic, 5023 ntl	1.5
3570	X56832	H. sapiens ENO3 gene for muscle specific enolase	0.21
3571	XM 011264	Homo sapiens KIAA0372 gene product (KIAA0372), mRNA	0
3572	NM 022628	Rattus norvegicus nephrin (Nphs1), mRNA	0.017
3573	AJ299718	Homo sapiens partial MASP2 gene, intron 7	6E-33
		Human DNA sequence from cosmid L75B9, Huntington's Disease	
	Z69651	Region, chromosome 4p16.3	0.022

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Streptococcus thermophilus Peb1 (peb1), GlnQ (glnQ), response	
		regulator Rr1 (rr1), Hpk2 (hpk2), YycJ-like protein (yycJ), and MurM	
3575	AF327739	(murM) genes, complete cds	0.59
		Drosophila melanogaster (P1 DS00397 (D18)) DNA sequence,	
3576	AC001656	complete sequence	0.068
		Periplaneta americana 16S ribosomal RNA gene, partial sequence;	
3577	AF262620	mitochondrial gene for mitochondrial product	0.2
		Zea mays auxin-binding protein (abp1) gene, exons 1-5 and complete	
3578	L08425	cds	0.64
3579	M97702	Drosophila melanogaster glutathione S-transferase gene	1.8
		Candidatus Carsonella ruddii natural-host Arytaina geniotae RNA	
		polymerase beta subunit (rpoB) and RNA polymerase beta-prime	
3580	AF268059	subunit (rpoC) genes, partial cds	0.063
2501	********	Entamoeba histolytica pyruvate: ferredoxin oxidoreductase (pfo1)	
3581	U30149	gene, complete cds	1.9
3582	X88893	C.jacchus intron 4 of visual pigment gene (green allele)	0.00008
3584	AK000939	Homo sapiens cDNA FLJ10077 fis, clone HEMBA1001864	0.19
2505	AT 114256	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen	
3585	AL114356	deprivation	5.6
3586	AF126992	Oxytropis besseyi var. ventosa chloroplast tRNA-Leu (trnL) gene,	
3587	Z47046	intron sequence Human cosmid QLL2C9 from Xq28	5.7
3307	24/040	Lactococcus lactis subsp. lactis IL1403 section 8 of 218 of the	3.7
3588	AE006246	complete genome	5.3
3300	AE000240	Oryza sativa microsatellite MRG1246 containing (AT)X23, genomic	3,3
3589	AY018921	sequence	0.71
3303	A1010921	Human DNA sequence from clone RP11-64M7 on chromosome 6,	0.71
3590	AL589705	complete sequence [Homo sapiens]	0.069
3370	AL307703	Mus musculus gamma-aminobutyric acid (GABA-A) receptor, subunit	0.005
3591	NM 010250	alpha 1 (Gabra1), mRNA	3.8
3592	AC024748	Caenorhabditis elegans cosmid Y110A2AR, complete sequence	0.017
3593	AJ002240	Mus musculus minisatellite tandem repeat (MMS9)	1.7
		Brassica napus gene for putative corticosteroid binding protein and	
3594	AJ293574	partial genc for hypothetical protein, cultivar N-o-9	0.21
		Homo sapiens myosin phosphatase target subunit 2 (MYPT2) gene,	
3596	AF324889	exon 1	0.22
		IGF2=insulin-like growth factor 2 {clone PPA1, exon 9} [human,	
3597	S62623	Genomic, 811 nt]	5.4
3598	Z22707	S.scrofa spl IFN gene	0.57
		Glycine max SG-05 gene for thiamin biosynthetic enzyme, complete	
3600	AB030490	cds	0.023
		CYP1A2=3-methylcholanthrene responsive gene [Oncorhynchus	
3602	S69278	mykiss=rainbow trout, liver, Genomic, 5023 nt]	1.7
3603	XM 029148	Homo sapiens hypothetical protein BC004923 (LOC85865), mRNA	0.002
3603	XM_029148	Homo sapiens hypothetical protein BC004923 (LOC85865), mRNA	0.002

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#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE L. concinnum chloroplast DNA for intergenic spacer between tRNA-Thr (UGU) and tRNA-Leu (UAA) 5'exon, collected from W. 3604 AJ002979 Pakawau, Cape Farewell, South Island, New Zealand 0.067 3605 U36927 Plasmodium yoelii rhoptry protein gene, complete cds 0.2 3606 AK021502 Homo sapiens cDNA FLJ11440 fis, clone HEMBA1001319 0.00001 Capra hircus Hsp70-3 gene for 70 kDa heat shock protein, complete AB033604 3607 0.18 3608 M81651 Human semenogelin II (SEMGII) gene, complete cds 1.8 3609 AB054512 Berardius bairdii DNA, SINE flanking sequence Tuti35 locus 0.49 Homo sapiens 17-beta-hydroxysteroid dehydrogenase IV (HSD17B4) 3610 AF057740 gene, exon 24 and complete cds 0.018 3611 AJ295158 Homo sapiens gene for thymosin beta-4 1E-48 3612 X79402 B.taurus mRNA for plasminogen 0.21 3613-NC 001807 Human mitochondrion, complete genome 3E-20 3614 X92204 P.hybrida NAM gene 0.049 Homo sapiens PDNP1 gene for phosphodiesterase I/nucleotide AB032016 pyrophosphatase 1, 5' flanking region and partial cds 0.009 3616 Macropus rufogriseus mannose-6-phosphate/insulin-like growth factor 3617 AF339159 II receptor (m6p/igf2r) mRNA, partial cds 0.47 0.0000001 3618 AB025285 Homo sapiens c-ERBB-2 gene, exons 1', 2', 3', 4' 3619 AJ002550 Homo sapiens MMP-1 gene, promoter region 3.8 3620 AB004856 Buchnera aphidicola mRNA for homoserine kinase, partial cds 0.79 3621 AB010426 Phytoplasma sp. gene for AL1 like protein, complete cds 1.8 Saccharomyces cerevisiae mitochondrion 21S ribosomal RNA (21S rRNA) gene, 21S R2 ribosomal RNA (21S R2 rRNA) gene, and ORF 3622 L36903 0.022 3623 AF267642 Filobasidiella neoformans var. neoformans Myo2p-like protein gene 0.49 XM 044499 Homo sapiens KIAA0484 protein (KIAA0484), mRNA 0.0001 3624 3625 L04550 Homo sapiens DNA fragment 2.6 Homo sapiens bone morphogenetic protein 7 (BMP7) gene, promoter region and partial cds 3626 AF210054 2.5 Human DNA sequence from clone RP11-67I12 on chromosome X. 3627 AL591505 complete sequence [Homo sapiens] 0.001 Homo sapiens mRNA; cDNA DKFZp564A072 (from clone 3628 AL049969 DKFZp564A072) 0.35 3629 AB055284 Macaca fascicularis brain cDNA, clone:OflA-11654 0.15 Homo sapiens vitamin D receptor-interacting protein complex 1.3 3630 AF304448 component DRIP150 (DRIP150) mRNA, complete cds 3631 AL133157 0.6 S.pombe chromosome I cosmid c1B2 Homo sapiens Chromosome 22q11.2 PCR Product dgcr-gapl In 0.005 3632 AC002475 DGCR Region, complete sequence Mus musculus 18 days embryo cDNA, RIKEN full-length enriched 3633 AK003943 library, clone:1110028F11, full insert sequence 4E-18 hemolin=48 kda C2-type immunoglobulin-like protein [Hyalophora 3634 S78712 cecropia=giant silkmoths, Genomic, 6622 nt] 3.5

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	-	Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUI
3635	AK023897	Homo sapiens cDNA FLJ13835 fis, clone THYRO1000712	2E-98
3636	X55978	T. brucei ESAG 8 gene for a leucine-rich repeat family protein	0.14
3637	XM_031534	Homo sapiens similar to uterine protein (M. musculus) (LOC90410), mRNA	5E-47
3638	AF044287	Drosophila melanogaster delta adaptin subunit of AP-3 (garnet) gene, complete cds	0.004
3639	AB060917	Macaca fascicularis brain cDNA clone:QtrA-14684, full insert sequence	1.1
3640	AK008522	Mus musculus adult male small intestine cDNA, RIKEN full-length enriched library, clone:2010305C02, full insert sequence	0.41
3641	NM_031340	Rattus norvegicus timeless (Drosophila) homolog (Timeless), mRNA	0.18
3642	AF334544	Homo sapiens chromosome Y landmark: proximal chromosome 1 translocation boundary distal to AZFc region	0.19
3643	XM_033642	Homo sapiens nucleophosmin (nucleolar phosphoprotein B23, numatrin) (NPM1), mRNA	1E-66
3645	AE001462	Helicobacter pylori, strain J99 section 23 of 132 of the complete genome	1.3
3646	L81761	Homo sapiens (subclone 6_c10 from P1 H17) DNA sequence, complete sequence	4E-12
3647	AJ271348	Homo sapiens partial DRD3 gene for dopamine D3 receptor, exon 1	0.19
		Simian immunodeficiency virus from African green monkey, tantalus	
3648	U58991	species (SIVtan) proviral DNA, complete genome	0.16
3649	NM_014601	Homo sapiens EH-domain containing 2 (EHD2), mRNA	0.087
3650	NM_032717		0.62
3651	XM_004462	Homo sapiens signal sequence receptor, alpha (translocon-associated protein alpha) (SSR1), mRNA	0,79
3652	AY016336	Phragmites australis rpl36-rps8 intergenic spacer, chloroplast sequence	0.69
3653	Z63374	H.sapiens CpG island DNA genomic Mse1 fragment, clone 82b3, forward read cpg82b3.ft1a	0.22
3654	U10927	Staphylococcus aureus M type 1 capsular polysaccharide biosynthesis (capA, capB, capC, capD, capE, capF, capG, capH, capI, capI, capK, capL, capM) genes, complete cds	0.026
3655	AE007446	Streptococcus pneumoniae section 129 of 194 of the complete genome	0.18
3656	BC006035	Mus musculus, clone IMAGE:3492938, mRNA, partial cds	5.3
3657	AF352740	Oncorhynchus mykiss clone OMM1055 microsatellite sequence	2.1
3658	NM_032141	Homo sapiens hypothetical protein DKFZp434K1421 (DKFZP434K1421), mRNA	0
		Phloeomyzus passerinii 12S small subunit ribosomal RNA gene, partial sequence; tRNA-Val gene, complete sequence; and 16S large	
3659	AF275231	subunit ribosomal RNA gene, partial sequence; mitochondrial genes for mitochondrial products	0.077

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		Tuble 374 Treatest Neighbor (Blash Va. Geneank)	
NO NO	ACCESSN	DESCRIP	D 1747 FTF
NO	ACCESSIV	Schizosaccharomyces pombe TFIIH subunit p47 (tfh47) gene,	P VALUE
3660	AF017646	complete cds	0.70
3661	AF200526	Zea mays cellulose synthase-2 (CesA-2) mRNA, complete cds	0.72
3001	AF200526	zea mays cenniose synthase-z (CesA-z) mkNA, complete cds	2.2
3662	AJ297654	Sus scrofa AACT2 gene for alpha-1-antichymotrypsin 2, exons 1-5	0.74
3663	XM_018189	Homo sapiens hypothetical protein FLJ13949 (FLJ13949), mRNA	0.74
3664	AE004047	Xylella fastidiosa 9a5c, section 193 of 229 of the complete genome	2.2
3665	XM_003012	Homo sapiens EphB1 (EPHB1), mRNA	6,9
3666	AF083240	Shewanella putrefaciens multi-cytochrome gene cluster, complete sequence; ferrous iron transporter (feoB), deca-heme c-type cytochrome (mtrO, -cype cytochrome precursor (mtrA), and outer membrane; protein precursor (mtrB) genes, complete cds	0.085
3667	AF100304	Caenorhabditis elegans cosmid W07B3, complete sequence	0.74
		Homo sapiens genomic DNA, chromosome 21q21.1-q21.2	
3668	AP001348	clone:f39G8, LL56-APP region, complete sequence	2E-18
3669	AB005039	Alternaria alternata MAT2 gene, complete cds	6.4
3670	NM_009181	Mus musculus sialyltransferase 8 (alpha-2, 8-sialytransferase) B (Siat8b), mRNA	0.22
3671	AB045894	Nepenthes alata NaAP4 mRNA for aspartic proteinase 4, complete cds	7
3672	M74944	Simian immunodeficiency virus (32H Re-isolate of SIVmac 251, clone W9) env gene, complete cds	0.25
3673	AF288211	Danio rerio Nkx5-1 mRNA, complete cds	0.072
3674	AC006753	Caenorhabditis elegans cosmid Y40A1A, complete sequence	0.02
3675	S53497	immunoglobulin epsilon chain constant region=secreted form {3' region} [human, B cell myeloma U-266, Genomic, 3198 nt]	6.8
		Human DNA sequence from clone RP11-85E24 on chromosome 6.	
3676	AL512305	complete sequence [Homo sapiens]	0.009
3677	AF207862	Bos taurus bradykinin receptor B2 (BDKRB2) gene, promoter	5.8
		Staphylococcus epidermidis strain SR1 clone step.1003e03 genomic	
3678	AF269417	sequence	5.1
3679	X53232	Rat mRNA for preoptic regulatory factor-2 (PORF-2)	2
3680	AB005039	Alternaria alternata MAT2 gene, complete cds	6.5
		Drosophila melanogaster genomic scaffold 142000013385369,	
3681	AE003252	complete sequence	1.9
3682	U19744	Caenorhabditis elegans integrin beta pat-3 gene, complete cds	0.083
3683	XM_010829	Homo sapiens potassium channel, subfamily K, member 3 (TASK-1) (KCNK3), mRNA	0.25
3684	AF065853	Homo sapiens OR7E11P pseudogene, partial sequence	0.077
3685	AF325859	Choristoneura fumiferana antifreeze protein (AFP-Lu1) gene, complete eds	2.3
3686		Homo sapiens breast cancer 2, early onset (BRCA2), mRNA	0,23

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WO0214500 [fla //E /WO0214500 opc]

#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEQ ID NO ACCESSN DESCRIP P VALUE 3687 AE002139 Ureaplasma urealyticum section 40 of 59 of the complete genome 0.17 3688 J00241 Human Ig germline kappa-L chain, C region (inv3 allele) Bos taurus von Willebrand factor gene, 5' flanking region and exon 1 3689 U23855 0.78 Mus musculus adult male testis cDNA, RIKEN full-length enriched 3690 AK015465 library, clone:4930455H04, full insert sequence 6.2 3691 XM 042056 Homo sapiens LOC86123 (LOC86123), mRNA Homo sapiens mRNA; cDNA DKFZp434E2221 (from clone 3692 AL137458 DKFZp434E2221) 0.086 3693 Z81079 Caenorhabditis elegans cosmid F39H11, complete sequence 2 Pteronotus parnellii NADH dehydrogenase subunit 1 (ND1) gene. 3694 AF142998 partial cds; mitochondrial gene for mitochondrial product 0.69 Mus musculus 11 days embryo cDNA, RIKEN full-length enriched 3695 AK012282 library, clone:2700023124, full insert sequence 0.75 Cucumis sativus CS-ACS1 gene for 1-aminocyclopropane-1-3696 AB032937 carboxylate synthase, complete cds 1.5 3697 M92295 Gorilla gorilla gamma-1 and gamma-2 globin genes, complete cds 0.029 AF034085 3698 Caenorhabditis elegans UNC-45 (unc-45) gene, complete cds 0.0001 Macaca fascicularis brain cDNA, clone:QnpA-14055 3699 AB048996 3700 AK024527 Homo sapiens cDNA: FLJ20874 fis, clone ADKA02818 0.000001 Mus musculus 14 days embryo liver cDNA, RIKEN full-length AK014489 3701 enriched library, clone:4432412D04, full insert sequence 4.3 3702 NM 022599 Rattus norvegicus outer membrane protein (Omp25), mRNA 1E-14 3703 AF177242 Bodo saltans clone pBME40 mitocondrial minicircle DNA 0.021 Homo sapiens clone NIGMS NA10926D chromosome 10 paralogous 3704 AF183329 5E-61 sequence variant, genomic sequence Saxifragella bicuspidata maturase K (matK) gene, partial cds: chloroplast gene for chloroplast product 3706 AF374728 0.65 Mus musculus adult male tongue cDNA, RIKEN full-length enriched 3707 AK010113 library, clone:2310068J16, full insert sequence 5E-29 3708 XM 045657 Homo sapiens McKusick-Kaufman syndrome (MKKS), mRNA 1.8 H. sapiens D2-1-K2G7np gene for immunoglobulin kappa chain 3709 AJ223690 variable region 0.026 3710 XM 008389 Homo sapiens Clu-related factor (CRF), mRNA 0.026 3711 AF200526 Zea mays cellulose synthase-2 (CesA-2) mRNA, complete eds 2.3 Mycoplasma pneumoniae M129 section 31 of 63 of the complete 3712 AE000049 genome 0.68 Homo sapiens cyclic nucleotide gated channel alpha 3 (CNGA3), XM 002495 3713 2.1 3714 AF322106 Oncorhynchus mykiss lysozyme II precursor, gene, complete cds Homo sapiens cDNA FLJ11145 fis, clone PLACE1006626, highly 3715 AK002007 0.029 similar to Homo sapiens mRNA for KIAA0928 protein Caenorhabditis elegans cosmid ZC302, complete sequence 3716 Z73978 Rattus norvegicus apoptotic protease activating factor 1 (Apaf1), 3717 NM 023979 mRNA 0.52

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#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE Oryza sativa hemoglobin 1 (hb1), hemoglobin 3 (hb3), and 3718 AF335504 hemoglobin 4 (hb4) genes, complete cds 0.077 Mus musculus, Similar to RIKEN cDNA 3230401N03 gene, clone MGC:6906 IMAGE:2655807, mRNA, complete cds 3719 BC006943 2 Mus musculus adult male testis cDNA, RIKEN full-length enriched 3720 AK007159 library, clone:1700110101, full insert sequence 6 3721 AF033002 Cerebratulus lacteus body wall globin gene, complete cds 3722 X95276 P.falciparum complete gene map of plastid-like DNA (IR-B) 0.2 Homo sapiens DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 8 3723 XM 012629 (RNA helicase) (DDX8), mRNA 1.4 Homo sapiens hypothetical protein DKFZp434G131 3724 (DKFZP434G131), mRNA XM 042643 0.69 Drosophila melanogaster clone GH07346 BcDNA.GH07346 3725 AF145640 (BcDNA.GH07346) mRNA, complete cds 2.3 Bovine COX7cP1 retroposed pseudogene for cytochrome c oxidase 3726 X58823 subunit VIIc 0,21 3727 XM 052301 Homo sapiens hypothetical protein MGC3199 (MGC3199), mRNA 6E-25 Dictyostelium mucoroides DMUC2 clone p288m2 G1-like and G5-3728 U00176 like ORFs' proteins, complete cds 0.00009 AJ132901 Drosophila guanche mitochondrial A+T-rich region 0.071 Human chromosome 14 DNA sequence Partial sequence from BAC R-1109N18 PCR1 of library RPCI-11 from chromosome 14 of Homo 3730 AL583762 sapiens (Human), complete sequence 7E-35 Mus musculus adult male corpus striatum cDNA, RIKEN full-length 3731 AK021083 enriched library, clone:C030015A19, full insert sequence XM_001934 Homo sapiens PRO0529 protein (PRO0529), mRNA 3732 1,4 AJ235836 Labordia tinifolia chloroplast ndhF gene 22 3734 M81855 Rat mdr mRNA sequence 1.9 Rattus norvegicus killer cell lectin-like receptor subfamily G, member 3735 NM 031649 1 (Klrg1), mRNA 49 3736 Caenorhabditis elegans cosmid F42E8, complete sequence Z78014 0.002 Homo sapiens chondroitin sulfate proteoglycan 6 (bamacan) (CSPG6). XM 045320 mRNA 3737 0.0005 AF288788 3738 Staphylococcus aureus rot-like protein Rlp (rlp) gene, complete cds 0.4 D16417 Dictyostelium discoideum mRNA 0.008 3740 X15465 Hamster SPC14 LINE L1 repeat DNA 0.000003 3741 Y14544 Danio rerio mRNA for Hoxe8 protein 0.014 Rattus norvegicus thyrotropin beta subunit (TSH-beta) gene, complete 3742 M13897 cds, clones RP100-14 and RP21 3.5 Plasmodium falciparum chromosome 2, section 27 of 73 of the 3743 AE001390 complete seguence 0.018 3744 XM 035389 Homo sapiens ribosomal protein, large, P1 (RPLP1), mRNA 9E-36 Homo sapiens alkylation repair: alkB homolog (ABH), mRNA 0.057

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	- DESCRIP	P VALUE
3746	AE007425	Streptococcus pneumoniae section 108 of 194 of the complete genome	0.15
3747	Z38113	S.cerevisiae chromosome IX lambda clone 4554	4.9
3749	X80491	S.latifolia SLM4 mRNA	0.02
3750	XM_035466	Homo sapiens hBKLF for basic kruppel like factor (LOC51274), mRNA	2E-35
3751	Z75957	Human DNA sequence from cosmid U203H4, between markers DXS366 and DXS87 on chromosome X	0.000001
3752	AE000800	Methanobacterium thermoautotrophicum from bases 58271 to 68710 (section 6 of 148) of the complete genome	0.19
3754	AJ409501	Mus musculus RNA binding site for Dazl protein, clone gf10	0.73
3755	AJ243829	Oryza sativa RPA gene for protein phosphatase 2A A subunit, exons 1- 12	3E-25
3756	AK026692	Homo sapiens cDNA: FLJ23039 fis, clone LNG02242	0.022
3757	S81000	{microsatellite regions} [Cryptococcus neoformans=pathogenic yeast, var. gattii, serotype B, UCLA 371-B, human isolate, Genomic, 283 nt]	0.94
3758	AF252420	Bos taurus clone MNB-5 microsatellite sequence	0.074
3759	AF338734	Homo sapiens hypothetical PHD zinc finger protein XAP135 pseudogene, complete sequence	2.1
3760	AJ401038	Human immunodeficiency virus type 1 proviral gp160 gene for envelope protein, strain 97DC.KFE267	0.79
3761	NC_002087	Plasmid pRL765, complete sequence	0.71
3762	AE004161	Vibrio cholerae chromosome I, section 69 of 251 of the complete chromosome	1.9
3763	Z54349	H.sapiens MN/CA9 GENE	0.56
3764	AF372396	HIV-1 isolate CM53658 from Cameroon envelope glycoprotein (env) gene, partial cds	1.8
3765	AY021127	Oryza sativa microsatellite MRG3452 containing (TA)X19, genomic sequence	1.9
3766	AK021485	Homo sapiens cDNA FLJ11423 fis, clone HEMBA1001024	3.4
3767	AF269468	Staphylococcus epidermidis strain SR1 clone step.1005a02 genomic sequence	0.69
3768	AF229080	Rattus norvegicus glucagon receptor gene, promoter and partial cds	0.71
3769	X04384	Sea urchin (L.pictus) gene for testis specific histone H2B-2	0.23
3770	Z36061	S. cerevisiae chromosome II reading frame ORF YBR192w	0.087
3771	AB038488	Halimeda discoidea chloroplast rbcL gene for ribulose-1,5- bisphosphate carboxylase/oxygenase large subunit, partial cds	0.25
3772	XM_003759	Homo sapiens leukocyte cell-derived chemotaxin 2 (LECT2), mRNA	0.000001
3773	AF122979	Tridacna maxima 16S ribosomal RNA gene, partial sequence; mitochondrial gene for mitochondrial product	0.076
3774	L44120	Homo sapiens (clone pHK1.5D) CMT1A gene, primer binding site	5E-39

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
3775	AC008855	Homo sapiens chromosome 5 clone CTD-2178M23, complete sequence	0.73
		Candidatus Carsonella ruddii natural-host Tainarys sordida ATP	
		synthase alpha subunit (atpA) gene, partial cds, ATP synthase gamma	
		subunit (atpG) gene, complete cds; and ATP synthase beta subunit	
3776	AF267225	(atpD) gene, partial cds	0.024
3777	U67231	Human clone HS4.74 Alu-Ya5 sequence	2E-11
		Homo sapiens similar to HYPOTHETICAL PROTEIN KIAA0379 (H.	
3778	XM_038985	sapiens) (LOC91526), mRNA	0.13
3779	X71625	H.sapiens dinucleotide repeat polymorphism cA14	0.057
		Mus musculus, clone MGC:7570 IMAGE:3493227, mRNA, complete	
3780	BC007480	cds	0.59
		Human immunodeficiency virus type 1 isolate ID4-77, clone 12, from	
3781	U60153	USA, envelope glycoprotein (env) gene, partial cds	0.013
		Homo sapiens hypothetical protein DKFZp564O1664	
3782	XM_050963	(DKFZP564O1664), mRNA	0
3783	AC024236	Homo sapiens BAC clone RP11-400I17 from Y, complete sequence	0.0001
3784	M17627	Chicken alpha-A-crystallin gene, complete cds and 5' flank	0.077
3785	NM_031557	Rattus norvegicus Prostaglandin I2 (prostacyclin) synthase (Ptgis), mRNA	5.8
3786	Z79647	B.thayeri mitochondrial 16S rRNA gene	0.23
3787	AF032386	Nicotiana tabacum aldose-1-epimerase-like protein (GP40) mRNA, complete cds	0.077
3788	X89568	P.sativum mRNA for HMGI/Y protein	5.8
3789	Z71180	Caenorhabditis elegans cosmid F22E12, complete sequence	0.24
3790	AJ239079	Tetrahymena pyriformis partial gc4 gene for guanylyl cyclase, isoform 4	0.79
		Dictyostelium discoideum PcmA (pcmA) and RacE (racE) genes.	
3791	AF3 10890	complete cds; tRNA-Phe gene, complete sequence; and unknown gene	0.00003
		Chilo iridescent virus type 6 zinc finger protein and non-histone	
		chromosomal high mobility group protein homolog genes, complete	
3792	L22300	cds	0.021
3793	AF155054	Rickettsia slovaca protein PS 120 (D) gene, partial cds	0.47
3794	X59799	M.musculus S-antigen gene promoter region	0.2
3795	AF330221	Oncorhynchus keta clone Oke4 microsatellite sequence	0.56
3796	XM 043227	Homo sapiens similar to proteasome (prosome, macropain) 26S subunit, non-ATPase, 13 (H. sapiens) (LOC92153), mRNA	6
3797	D17668	Clostridium septicum gene for alpha-toxin, complete cds	0.39
		Mycoplasma mycoides mycoides SC strain Afade hypothetical proline rich glycoprotein, hypothetical surface-located membrane protein, lipoprotein B precursor (lipB), and hypothetical ABC transporter	
3798		protein genes, complete cds; hypothetical ABC transport>	1.5

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Rhagigaster sp. M222 16S ribosomal RNA gene, partial sequence:	
3799	AF379892	mitochondrial gene for mitochondrial product	0.068
3800	ND 6 004050		
3801	NM_024379 AF364550	Rattus norvegicus glutamate receptor delta-2 subunit (Grid2), mRNA  Drosophila melanogaster transposon cruiser, complete sequence	1.5 5.4
3601	AI-304550	Diosophila melanogaster transposon cruiser, complete sequence	5,4
3803	AJ315160	Xenopus tropicalis BMP-2 gene for bone morphogenetic protein 2	0.24
3804	AJ293015	Pisum sativum lox1:Ps:7 gene for lipoxygenase, exons 1-8	0.079
3805	Z82768	R.prowazekii genomic DNA fragment (clone A810F)	1.5
3806	NM_013780	Mus musculus neuronal PAS domain protein 3 (Npas3), mRNA	0.72
3807	U56088	Human periodic tryptophan protein 2 (PWP2) gene, exons 3 to 14	0.25
3808	D88987	Mus musculus Ampd3 gene, exon 4	0.18
3809-	Z72680	S.cerevisiae chromosome VII reading frame ORF YGL158w	0.73
3810	M17988	Spiroplasma virus 4 (SpV4) replicative form, complete genome	5.3
3811	XM_039684	Homo sapiens putative G-protein coupled receptor (SH120), mRNA	0.17
	70		
3812	XM_039684	Homo sapiens putative G-protein coupled receptor (SH120), mRNA	0.19
3813	X60735	S.murinus Dq52 gene and immunoglobulin heavy chain S region gene loci	
3814			1.4 5E-14
3014	XM_045862	Homo sapiens ring finger protein 10 (RNF10), mRNA Homo sapiens ppar gamma gene for peroxisome proliferator activated-	3E-14
3815	AB005521	receptor gamma, exon 1	
3816	AF271964	Bos taurus microsatellite MNB-162 sequence	1.6 0.019
3817	AF271304 AF274307	Zaocys dhumnades creatine kinase mRNA, complete cds	6.1
3017	AL-274307	Mus musculus, clone MGC:6551 IMAGE:2655861, mRNA, complete	0.1
3818	BC010716	cds	5.5
3010	DC010710	Vibrio cholerae chromosome I, section 127 of 251 of the complete	3,3
3819	AE004219	chromosome	5.7
		Macaca fascicularis brain cDNA clone: OccE-22277, full insert	5.7
3820	AB060193	sequence	1.9
3821	AY007106	Homo sapiens clone TCCCIA00427 mRNA sequence	3E-45
3822	X02801	Mouse gene for glial fibrillary acidic protein (GFAP)	0.08
3824	AJ234506	Hordeum vulgare genomic DNA fragment; clone MWG0561.uni	2.1
		Mus musculus 11 days pregnant adult female ovary and uterus cDNA.	
		RIKEN full-length enriched library, clone:5031405K23, full insert	
3825	AK019864	sequence	1.8
3826	AK023304	Homo sapiens cDNA FLJ13242 fis, clone OVARC1000578	6,5
		Homo sapiens p53DINP1 gene for p53DINP1a, p53DINP1b, complete	
3827	AB062056	cds, alternative splicing	0.055
		Dendrocerus carpenteri 16S ribosomal RNA gene, partial sequence;	
3828	AF379858	mitochondrial gene for mitochondrial product	0.07
3829	X05621	Chironomus pallidivittatus BR1 gene for giant secretory protein	0.077

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		)	
NO	ACCESSN	DESCRIP	P VALUE
		Homo sapiens human endogenous retrovirus HERV-H10 pol protein	1 TILLOL
		(pol) gene, partial cds; env pseudogene and 3' LTR, complete	1
3830	AF108841	sequence	0.009
		Ixodes scapularis haplotype SSCP NC2_29 16S ribosomal RNA gene,	
3831	AF309030	partial sequence; mitochondrial gene for mitochondrial product	0.21
		Candidatus Carsonella ruddii natural-host Glycaspis brimblecombei	
2022	A TO COO 5 5	RNA polymerase beta subunit (rpoB) and RNA polymerase beta-prime	
3832	AF268055	subunit (rpoC) genes, partial cds	0,0007
3833	AY021307	Oryza sativa microsatellite MRG3632 containing (TA)X23, genomic sequence	0.000
3633	A1021307	Homo sapiens dual-specificity tyrosine-(Y)-phosphorylation regulated	0.009
3834	XM 034551	kinase 4 (DYRK4), mRNA	0,00003
5051	7114_051551	Homo sapiens cDNA FLJ10785 fis, clone NT2RP4000457, weakly	0.00003
		similar to UBIQUITIN CARBOXYL-TERMINAL HYDROLASE 15	
3835	AK001647	(EC 3.1.2.15)	۱ ،
3836	XM_043492	Homo sapiens 42259 (KIAA1728), mRNA	5E-19
3837	U22345	Human chromosome 20q12 locus-specific repeat	0.003
3838	J05535	Bovine vascular smooth muscle connexin43 mRNA, complete cds	0.18
3839	AF290221	Cucumis sativus clone B80F mitochondrial genomic sequence	0.065
3840	NM_010059	Mus musculus disrupted meiotic cDNA 1 homolog (Dmc1h), mRNA	5.3
3841	NM_017681	Homo sapiens hypothetical protein FLJ20130 (FLJ20130), mRNA	2
3842	AE002114	Ureaplasma urealyticum section 15 of 59 of the complete genome	1.9
3843	AK022372	Homo sapiens cDNA FLJ12310 fis, clone MAMMA1001970	0.019
2044	1122622		
3844	U33633	Rice ragged stunt virus minor structural protein gene, complete cds	0.21
3845	AB062993	Macaca fascicularis brain cDNA clone:QmoA-10825, full insert sequence	4E-40
3043	ABOOZJIJ	sociation	4E-40
		Gorilla gorilla partial Imp7 gene for large multifunctional protease 7,	
3846	AJ251485	exon 7 and tap2 gene for antigen peptide transporter 2, exons 1-6	0.71
		g-p-p	0172
		Homo sapiens genomic DNA, chromosome 21q21.2, LL56-APP	
3847	AP000145	region, clone B2291C14-R44F3, segment 10/10, complete sequence	0.68
		Homo sapiens similar to ribosomal protein S3A (H. sapiens)	
3848	XM_046052	(LOC65465), mRNA	0.017
3849	XM_012036	Homo sapiens potassium inwardly-rectifying channel, subfamily J, member 5 (KCNJ5), mRNA	5
		Pseudoregma bambucicola mitochondrial genes for small subunit	
		rRNA, tRNA-Val, large subunit rRNA, partial and complete	
3850	AB035883	sequences, country:Taiwan:Sun Moon Lake	0.077
3851	AB037688	Xenopus laevis Xptch-2 mRNA for patched-2, complete cds	1.8

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALU
3852	AB041264	Mus musculus pseudogene for GTP-binding protein beta1 subunit	0.53
		Plasmodium falciparum isolate MM cytoadherence linked asexual	
3853	AF288170	protein (clag9) mRNA, partial cds	0.083
		Neospora caninum RNA polymerase C1 (rpoC1) gene, partial	
3854	AF304321	sequence, plastid gene for plastid product	0.002
3855	45100101	Peziza varia 18S ribosomal RNA gene, and internal transcribed spacer	
3633	AF133181	1, partial sequence	2.1
3856	NM 012694	Rattus norvegicus Solute carrier family 6 (neurotransmitter	
3630	1414_012094	transporter, dopamine), member 3 (Slc6a3), mRNA	0.56
3857	NM 030261	Mus musculus hypothetical protein MGC7182 (MGC7182), mRNA	6E-36
5001	1412_030201	Pseudosymblepharis schimperiana tRNA-Leu, partial sequence; trnL-	0E-30
		trnF intergenic spacer, complete sequence; and tRNA-Phe, partial	
3858	AF231164	sequence; chloroplast genes for chloroplast products	0.059
3859	Z36070	S.cerevisiae chromosome II reading frame ORF YBR201w	0.064
		H. sapiens (D1S489) DNA segment containing (CA) repeat; clone	0.004
3860	Z24279	AFM309ve9; single read	0.2
3861	XM_036937	Homo sapiens KIAA1020 protein (KIAA1020), mRNA	2.1
3862	AB033019	Homo sapiens mRNA for KIAA1193 protein, partial cds	1.6
		Homo sapiens hypothetical protein DKFZp434D0513	
3864	XM_030075	(DKFZp434D0513), mRNA	2E-74
		Rattus norvegicus apoptotic protease activating factor 1 (Apaf1),	
3865	NM_023979	mRNA	0.59
3866	U50715	Mus musculus alpha-galactosidase A gene, complete cds	0.008
3867	AF078780	Caenorhabditis elegans cosmid C04F2	1.7
2050			
3869	L07305	Histoplasma capsulatum (clone pMS3) H-ATPase gene, complete cds	0.46
3870	XM 035796	Homo sapiens hypothetical gene supported by AF001893; AH005586	
36/0	AM_033796	(LOC91063), mRNA	1.7
3871	NM_025788	Mus musculus RIKEN cDNA 4930511N13 gene (4930511N13Rik), mRNA	
3872	AF339787	Homo sapiens clone IMAGE:205688, mRNA sequence	5,4
38/2	AF339767	Rattus norvegicus glial fibrillary acidic proteins alpha and delta	2
3873	AF028784	(GFAP) gene, alternatively spliced products, complete eds	0.005
3073	711 020704	Mus musculus lysyl-tRNA synthetase (Kars) gene, exon 14, complete	0.003
		cds; adenosine deaminase tRNA-specific 1 (Adat1) gene, exons EA, 1	
3874	AF328904	and partial cds	0.22
		Rhytidiadelphus triquetrus 23S ribosomal RNA gene, partial	0,22
		sequence; internal transcribed spacer 2, 4.5S ribosomal RNA gene,	
		internal transcribed spacer 3, and 5S ribosomal RNA gene, complete	
3875	AF033647	sequence, chloroplast genes for chloroplast RNAs	2
3876	BC006035	Mus musculus, clone IMAGE:3492938, mRNA, partial cds	5.6
3877		Homo sapiens spectrin beta IV (SPTBN3) mRNA, complete cds,	
		alternatively spliced	0.5
		Anguilla japonica GnRH receptor mRNA for gonadotropin-releasing	
3878	AB041327	hormone receptor, complete cds	0.24

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	-	Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		gas ( and standard)	
NO	ACCESSN	DESCRIP	P VALUE
		Homo sapiens acid ceramidase (ASAH) gene, exons 5 through 14, and	1 TIEGE
3879	AF220175	complete cds	0.0009
3880	X89398	H.sapiens ung gene for uracil DNA-glycosylase	0.002
		Staphylococcus aureus methicillin-resistance protein (mecR) gene and	
3881	L14017	unknown ORF, complete cds	0.56
3882	M97008	Xenopus laevis/gilli hybrid IgM chain gene, clone LG7Gmu	0.007
3883	Z78414	Caenorhabditis elegans cosmid W09D12, complete sequence	0.63
3884	AL512549	S.pombe chromosome I BAC pB2B4	1.9
		Mus musculus Chetk-alpha gene for choline/ethanolamine kinase-	
3885	AB030616	alpha, exon 1 and 5'-flanking region	6.4
		Bradypus tridactylus 16S ribosomal RNA gene, mitochondrial gene	
3886	AF069535	for mitochondrial RNA, complete sequence	1.5
3887	AK023304	Homo sapiens cDNA FLJ13242 fis, clone OVARC1000578	0.056
3888	4 Donogra	Brassica rapa SP11-32 mRNA for S locus protein 11-32, partial cds,	
3888	AB039756	strain:S32(50-11t)	1.9
3889	AL512405	Human DNA sequence from clone RP11-370N21 on chromosome 6,	
3890	AF147813	complete sequence [Homo sapiens] Protrama radicis elongation factor 1 alpha gene, partial cds	1.6
3890	AF14/013	Troutanta radicis elongation factor 1 alpha gene, partial cds	0.18
		Homo sapiens region containing variable charge protein on X with eight repeats; variable charge protein on X with two repeats; variable	
3891	XM 046967	charge, X chromosome (LOC92738), mRNA	0.01
3071	2111_040907	Charge, A chiomosome (EOC52738), mrdvA	0.21
3892	XM 037664	Homo sapiens hypothetical protein MGC4090 (MGC4090), mRNA	1.5
		Silene foetida partial rpb2 gene for RNA polymerase II, exons 23 and	1.5
3893	AJ296142	24	0.16
		Arabidopsis thaliana putative pectinesterase (K19E1.17/AT5g53370)	
3894	AF360340	mRNA, complete cds	2.7
		Hentzia palmarum 16S ribosomal RNA, partial sequence;	
3895	AF327975	mitochondrial gene for mitochondrial product	0.55
3896	Z15140	L.esculentum mRNA for chitinase	2.1
1		Chlamydomonas reinhardtii chloroplast complete Eco10 fragment,	
3897	U57326	rpoC2 gene, partial cds	0.001
3898	AK024908	Homo sapiens cDNA: FLJ21255 fis, clone COL01321	0.003
3899	NC_002087	Plasmid pRL765, complete sequence	0.62
3900	AF063220	Papaya ringspot virus isolate P polyprotein gene, partial cds	0.25
2001	15000150	Chlamydophila pneumoniae AR39, section 9 of 94 of the complete	
3901	AE002173	genome	0.25
3902	BC001152	Homo sapiens, growth arrest-specific 7, clone MGC:1347	
3902	AB011800	IMAGE:3353809, mRNA, complete cds	5.9
3903	ABUITAUU	Agrobacterium tumefaciens plasmid pRiA4 gene, virB operon	5.4
3904	BC010389	Homo sapiens, clone MGC:13615 IMAGE:4283814, mRNA, complete cds	
3304	DC010369	Oncorhynchus mykiss non-classical MHC class I antigen (Onmy-	0.084
3905	AF091779	UAA) gene, Onmy-UAA*0101 allele, complete cds	2.3
3703		Plasmodium falciparum clone 3D7 unconventional myosin PfM-B	2.3
3906	AF222716	gene, complete cds	6.2
2700		Barre, compress ods	0.2

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN		
3907	Z48951	DESCRIP S.cerevisiae chromosome XVI cosmid 9723	P VALUE
3907	XM 040326		2.5
3909	AF271964	Homo sapiens absent in melanoma 1 (AIM1), mRNA	0.056
3909	AF2/1964	Bos taurus microsatellite MNB-162 sequence	0.019
3910	BC003261	Mus musculus, scrine/threonine kinase 5, clone MGC:5803 IMAGE:3501444, mRNA, complete cds	
3911	AJ011641	Arabidopsis thaliana (ecotype Columbia) spl8 gene, exons 1-3	0.056
3912	AK000864	Homo sapiens cDNA FLJ10002 fis, clone HEMBA1000046	1.7
3913	XM 045747		0.53
3913	AW_043747	Homo sapiens midline 1 (Opitz/BBB syndrome) (MID1), mRNA	2E-33
3914	AF269581	Staphylococcus epidermidis strain SR1 clone step.1011g07 genomic sequence	
3714	AF209381	sequeixe	0.057
3915	XM_042333	Homo sapiens casein kinase 2, beta polypeptide (CSNK2B), mRNA	1.9
		Staphylococcus epidermidis strain SR1 clone step.1023h06 genomic	
3916	AF269810	sequence	0.022
3917	AJ404228	Saccharomyces douglasii mitochondrial tRNA gene cluster	0.21
		Homo sapiens hypothetical gene supported by AK021954	
3918	XM_027223	(LOC89916), mRNA	0.61
3919	AJ011641	Arabidopsis thaliana (ecotype Columbia) spl8 gene, exons 1-3	2
ĺ		Gallus gallus unoccupied integration site for endogenous ALV-type	
3920	AF020725	retroviral element ev-B2	2
		Human DNA sequence from clone RP11-518C21 on chromosome	
3921	AL157825	Xq13.2-21.1, complete sequence [Homo sapiens]	1,2
3922	AF203676	Myotis myotis microsatellite H23 sequence	0.2
3923	XM_010734	Homo sapiens smg GDS-ASSOCIATED PROTEIN (Kifap3), mRNA	0.66
		Heterodera glycines beta-1,4-endoglucanase-3 precursor (eng-3) gene,	
3924	AF056048	promoter and complete cds	0.026
3925	XM 042676	Homo sapiens hypothetical protein MGC4618 (MGC4618), mRNA	2.2
		Human DNA sequence from clone RP11-24A10 on chromosome 13.	2.2
3926	AL359750	complete sequence [Homo sapiens]	0.024
	122337130	complete sequence [riolite suprens]	0.024
3927	NM 019054	Homo sapiens hypothetical protein MGC5560 (MGC5560), mRNA	2E-31
3928	AL023845	Caenorhabditis elegans cosmid Y51B9A, complete sequence	0.25
		Homo sapiens, ribosomal protein L44, clone MGC:2064	0.23
3929	BC001781	IMAGE:3353669, mRNA, complete cds	7E-43
		Mus musculus adult male lung cDNA, RIKEN full-length enriched	75.43
3930	AK004553	library, clone:1200002O22, full insert sequence	0.51
3932	X17339	Dengue-2 virus NS1 gene for nonstructural protein (patient M2)	0.23
		Vibrio cholerae chromosome I, section 186 of 251 of the complete	
3933	AE004278	chromosome	0.35
		Drosophila melanogaster putative extracellular ligand trunk gene,	0.55
3934		complete cds	0.53
		Homo sapiens alpha2B-adrenergic receptor (alpha2C2AR) gene.	0.55
3935		complete cds	5.2
3936		Homo sapiens cDNA: FLJ20946 fis, clone ADSE01819	5
		Atomic Suprems dorate A 202007-10 Ins, Citotic AD 0120/1019	

WC0214500 [fle://E/WO0214500 opc]

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SEQ_ID   NO   ACCESSN   DESCRIP	P VALUE 0.24 1.8 0.55 1.8 e-107 0.00002 0.64 0.71 1.9
3937   K00489   human enkephalin gene, intron c (3' end) and 3' flank	0.24 1.8 0.55 1.8 e-107 0.00002 0.64
3939   U22545   Gallus gallus tubby-like protein (TULP1) mRNA, complete ods   Human DNA sequence from cosmid L60G9B, Huntington's Disease   Region, chromosoms 4p16.5 contains EST\$   Dictyostclium discondeum random slug cDNA22 protein (rsc22) mRNA, complete ods   mRNA, complete ods   mRNA, complete ods   mRNA, complete ods   mRNA, complete ods   mRNA, complete ods   mRNA, complete ods   mRNA   Market   mRNA   mRNA   mRNA   Homo sapiens Na+,K+-ATPase gamma-subunit (FXYD2) gene, alternatively spliced isoforms 1 and 2, exons 1 through 6 and   complete ods   Homo sapiens Na+,K+-ATPase gamma-subunit (FXYD2) gene, alternatively spliced isoforms 1 and 2, exons 1 through 6 and   complete ods   Homo sapiens partial FAH gene for fumarylacatoacetate hydrolase, intron 10   introduced   market	1.8 0.55 1.8 e-107 0.00002 0.64 0.71
Human DNA sequence from cosmid L60G9B, Huntington's Disease Region, chromosome 4p16.3 contains ESTs Dictyostedium discoideum random slug cDNA22 protein (rsc22) mRNA, complete cds Miss musculus RIKEN cDNA 3110040N11 gene (3110040N11Rik), mRNA Horno sapiens RiKEN cDNA 3110040N11 gene (3110040N11Rik), mRNA Horno sapiens Na+,K+-ATPase gamma-subunit (f?XYD2) gene, alternatively spliced isoforms 1 and 2, exons 1 through 6 and complete cds Human 3-beta-hydroxysteroid dehydrogenase/delita-5-delita-4-isomerase (3-beta-HSD) gene, complete cds Horno sapiens partial FAH gene for fumarylacatoacetate hydrolase, intron 10 aspiens partial FAH gene for fumarylacatoacetate hydrolase, intron 10 aspiens partial FAH gene for fumarylacatoacetate hydrolase, intron 10 aspiens partial FAH gene for fumarylacatoacetate hydrolase, intron 10 aspiens partial FAH gene for fumarylacatoacetate hydrolase, intron 10 aspiens partial FAH gene for fumarylacatoacetate hydrolase, intron 10 aspiens gene for mydroli niligodentrocyte glycoprotein (MOG) asphylococcus epidermidis strain SR1 clone step. 1023b09 genomic sequence Chlosyne janais 16S ribosomal RNA gene, partial sequence; mitochondrial gene for mitochondrial product Hyoscyamus niger tr1 gene for tropinone reductase-I, complete cds Hyoscyamus niger tr1 gene for tropinone reductase-I, complete cds Homo sapiens profiba gene, intron 1, partial sequence; Homo sapiens profiba gene, intron 1, partial sequence; Homo sapiens profiba gene, intron 1, partial sequence; Homo sapiens profiba gene, intron 1, partial sequence; Homo sapiens profiba gene, intron 1, partial sequence; Homo sapiens profiba gene, intron 1, partial sequence; Homo sapiens profiba gene, intron 1, partial sequence; Homo sapiens profiba gene, intron 1, partial sequence; Homo sapiens profiba gene, intron 1, partial sequence; Homo sapiens profiba gene for mitochondrial product	0.55 1.8 e-107 0.00002 0.64
3930   Z69363   Region, chromosome 4p16.3 contains ESTS	1.8 e-107 0.00002 0.64 0.71
Dictyostelium discoideum random slug cDNA22 protein (rsc22)   mRNA, complete cds	1.8 e-107 0.00002 0.64 0.71
3941 U32516   mRNA, complete cds   Mins musculus RIKEN cDNA 3110040N11 gene (3110040N11Rik), mRNA   Homo sapiens Na+,K+-ATPase gamma-submit (PXYD2) gene, alternatively spliced isoforms 1 and 2, exons 1 through 6 and complete cds   Human 3-beta-hydroxysteroid dehydrogenase/delta-5-delta-4-isomerase (3-beta-HSD) gene, complete cds   Homo sapiens partial FAH gene for fumarylacatoacetate hydrolase, intro 10   3946 AB042193   Mus musculus mac25 gene promoter and exon 1, partial cds   April 27802500   Cryptosporidium parvum Hsp60 gene, partial cds   Cryptosporidium parvum Hsp60 gene, partial cds   April 27802500   Staphylococcus epidermidis strain SR1 clone step. 1023b09 genomic sequence   AF269768   AR022181   Homo sapiens CDNA FLJ12119 fis, clone MAMMA1000092   Chlosyne jamais 16S ribosomal RNA gene, partial sequence; mitochondrial gene for mitochondrial product   AF186889   Hyoscyamus niger tr1 gene for tropinone reductase-I, complete cds   4B026544   Hyoscyamus niger tr1 gene for tropinone reductase-I, complete cds   4B03879   Homo sapiens profiba gene, intron I, partial sequence;   Homo sapiens profiba gene, intron I, partial sequence   Homo sapiens profiba gene, intron I, partial sequence   Homo sapiens profiba gene, intron I, partial sequence   Homo sapiens profiba gene, intron II, partial sequence   Homo sapiens profiba gene, intron II, partial sequence   Legionetic falloni macrophage infectivity potentiator (migh) gene, (might profiba gene, intron II, partial sequence)   Homo sapiens profiba gene, intron II, partial sequence   Legionetic falloni macrophage infectivity potentiator (migh) gene, (might profiba gene, intron II)   Legionetic des   Legionetic falloni macrophage infectivity potentiator (might gene, partial sequence)   Legionetic falloni macrophage infectivity potentiator (might gene, partial sequence)   Legionetic falloni macrophage infectivity potentiator (might gene, partial sequence)   Legionetic falloni macrophage infectivity potentiator (might gene, partial sequence)   Legionetic falloni mac	e-107 0.00002 0.64 0.71
Mis musculus RIKEN cDNA 3110040N11 gene (3110040N11Rik), mRNA  Image: MRNA and the most spices Na+,K+-ATPase gamma-subunit (PXYD2) gene, alternatively spliced isoforms 1 and 2, exons 1 through 6 and complete cds  AF241235 complete cds  Human 3-beta-hydroxysteroid dehydrogenase/delta-5-delta-4-isomerase (3-beta-HSD) gene, complete cds  Human 3-beta-hydroxysteroid dehydrogenase/delta-5-delta-4-isomerase (3-beta-HSD) gene, complete cds  Homo spices partial FAH gene for fumarylacatoacetate hydrolase, intron 10  3945 AJ275020  3946 AB042198 Mis musculus mac25 gene promoter and exon 1, partial cds  Cryptosporidium parvum Hsp60 gene, partial cds  Cryptosporidium parvum Hsp60 gene, partial cds  Staphylococcus epidermidis strain SRI clone step 10/33609 genomic sequence  3949 AF269768  3950 AK02131 Homo saptens cDNA FLJ12119 fis, clone MAMMA 1000092  Chlosyne janais 16S ribosomal RNA gene, partial sequence; mitochondrial gene for mitochondrial product  M3266 Homo sapiens promoter tropinone reductase-1, complete cds  Hyoscyamus niger tr1 gene for tropinone reductase-1, complete cds  M3951 M12666 Homo sapiens FIGBs gene, intron 1, partial sequence  3955 Z72888 S. cerevisiae chromosome VII reading frame ORF YGR103w  Legionetti Ralonii macrophage infectivity potentiator (rim) gene, (rim) percophage infectivity potentiator (rim) gene,	e-107 0.00002 0.64 0.71
3942   NM_026077   mRNA	0.00002 0.64 0.71
Homo sapiens Na+,K+-ATPase gamma-subunit (FXYD2) gene, alternatively spliced isoforms 1 and 2, exons 1 through 6 and complete cds  Human 3-bda-hydroxysteroid dehydrogenase/delta-5-delta-4- isomerase (3-bda-HSD) gene, complete cds  Homo sapiens partial FAH gene for fumarylacatoacetate hydrolase, intron 10  3946 AB042198 Mus musculus mac25 gene promoter and exon 1, partial cds  AJ275020 Intron 10  3947 AF082520 Cryptosporidium parvum Hsq60 gene, partial cds  Cryptosporidium parvum Hsq60 gene, partial cds  AB042198 Mus musculus mac25 gene promoter and exon 1, partial cds  Cryptosporidium parvum Hsq60 gene, partial cds  AF082520 Cryptosporidium parvum Hsq60 gene, partial cds  Cryptosporidium parvum Hsq60 gene, partial cds  Cryptosporidium parvum Hsq60 gene, partial cds  Cryptosporidium parvum Hsq60 gene, partial cds  Cryptosporidium parvum Hsq60 gene, partial cds  Cryptosporidium parvum Hsq60 gene, partial cds  Cryptosporidium parvum Hsq60 gene, partial sequence; mitochondrial gene for mitochondrial product  Chlosyne janais 16S ribosomal RNA gene, partial sequence; mitochondrial gene for mitochondrial product  Hyoscyamus niger trl gene for tropinone reductase-I, complete cds  Hyoscyamus niger trl gene for tropinone reductase-I, complete cds  Homo sapiens gene for caspase-10, exon 11 and complete cds  Legionella fallonii macrophage infectivity potentiator (mig) gene, complete cds	0.00002 0.64 0.71
alternatively spliced isoforms 1 and 2, exons 1 through 6 and complete cds  3944 M38180 Human 3-beta-hydroxysteroid dehydrogenase/delta-5-delta-4-isomerase (3-beta-HSD) gene, complete cds  Human 3-beta-hydroxysteroid dehydrogenase/delta-5-delta-4-isomerase (3-beta-HSD) gene, complete cds  Human 3-beta-hydroxysteroid dehydrogenase/delta-5-delta-4-isomerase (3-beta-HSD) gene, complete cds  Human 3-beta-hydroxysteroid dehydrogenase/delta-5-delta-4-isomerase (3-beta-HSD) gene, complete cds  Human 3-beta-hydroxysteroid dehydrogenase/delta-5-delta-4-isomerase (3-beta-HSD) gene, complete cds  1946 AP042198 Missusculus mac25 gene promoter and exon 1, partial cds  Cryptosporidium parvum Hsp60 gene, partial cds  Cryptosporidium parvum Hsp60 gene, partial cds  Staphylococcus epidermidis strain SRI clone step.1023b09 genomic  Staphylococcus epidermidis strain SRI clone step.1023b09 genomic  3-940 AF046768  Staphylococcus epidermidis strain SRI clone step.1023b09 genomic  Staphylococcus epidermidis strain SRI clone step.1023b09 genomic  Chlosyne janais 16S ribosomal RNA gene, partial sequence;  mitochondrial gene for mitochondrial product  M32664 Homo sepiens TIGB3 gene, intron 1, partial sequence  3-955 AF04686 Homo sepiens TIGB3 gene, intron 1, partial sequence  3-955 AF04686 Homo sepiens TIGB3 gene, intron 1, partial sequence  3-956 AF046878 S. cerevisiae chromosome VII reading frame ORF YGR103w  Homo sepiens gene for caspase-10, exon 11 and complete cds  Legionella fallonii macrophage infectivity potentiator (mit) gene, fring partial sequence	0.64
3944	0.64
Human 3-beta-hydroxysteroid dehydrogenase/delta-5-delta-4- isomeras (2-beta-HSD) gene, complete cds Homo sspiens partial FAH gene for fumarylacatoacetate hydrolase, intron 10 3946 AB042198 Mus musculus mac25 gene promoter and exon 1, partial cds 1947 AF082520 Cryptosporidium parvum Hsp60 gene, partial cds 1948 Z48051 H. suplens gene for myelin oligodendrocyte glycoprotein (MOG) Staphylococcus cpidermidis strain SRI clone step 1023b09 genomic sequence 1949 AF269768 Saphylococcus cpidermidis strain SRI clone step 1023b09 genomic sequence 1951 AF186889 Chlosyne janais 16S ribosomal RNA gene, partial sequence, 1952 AB026544 Hyoscyamus niger tr1 gene for tropinone reductase-1, complete cds 1955 Z72888 S. cerevisiae chromosome VII reading frame ORF YGR103w 1950 AB038979 Homo sapiens gene for caspase-10, exon 11 and complete cds 1955 Legionelfa fallonii macrophage infectivity potentiator (migh) gene,	0.64
3944   M38180   isomerase (3-beta-HSD) gene, complete cds	0.71
Homo Sapiens partial FAH gene for fumarylacatoacetate hydrolase, intron 10	0.71
3945   A1275020   intron 10	
3947 AF082520 Cryptosporidium parvum Hsp60 gene, partial cds	
3947 AF082520 Cryptosporidium parvum Hsp60 gene, partial cds	
3948   Z48051   H. sapiens gene for myelin oligodendrocyte glycoprotein (MOG)	0.68
Staphylococcus epidermidis strain SR1 clone step. 1023b09 genomic sequence	0.71
3950 AK022181 Homo sapiens cDNA FLJ12119 fis, clone MAMMA1000092   Chlosyne jamais 16S ribosomal RNA gene, partial sequence;   3951 AF186889 mitochondrial gene for mitochondrial product   3952 AB026544 Hyoscyamus niger tr1 gene for tropinone reductase-I, complete cds   3954 MJ2666 Homo sapiens TrGB3 gene, intron I, partial sequence   3955 Z72888   S.cerevisiae chromosome VII reading frame ORF YGR103w     3956 AB038979 Homo sapiens gene for caspase-IO, exon II and complete cds   Legionella fallonii macrophage infectivity potentiator (mip) gene,	
AF186889 Chlosyne janais 16S ribosomal RNA gene, partial sequence; mitochondrial gene for mitochondrial product 3952 AB026544 Hyoscyamus niger tr1 gene for tropinone reductase-I, complete cds 3954 M12666 Homo sapiens ITGB3 gene, intron I, partial sequence 3955 Z72888 S. cerevisiae chromosome VII reading frame ORF YGR103w 4B038979 Homo sapiens gene for caspase-IO, exon I1 and complete cds Legionella fallonii macrophage infectivity potentiator (migh) gene,	0.59
3951         AF186889         mitochondrial gene for mitochondrial product           3952         AB026544         Hyoscyamus niger tr1 gene for tropinone reductase-I, complete cds           3954         M32666         Homo sapiens TTGB3 gene, intron 1, partial sequence           3955         Z72888         S.cerevisiae chromosome VII reading frame ORF YGR103w           3956         AB038979         Homo sapiens gene for caspase-10, exon 11 and complete cds           Legionella fallonii macrophage infectivity potentiator (mip) gene,	0.62
3952 AB026544   Hyoscyamus niger tr1 gene for tropinone reductase-I, complete cds   3954 M32666   Homo sapiens ITGB3 gene, intron I, partial sequence   3955 Z72888   S.cerevisiae chromosome VII reading frame ORF YGR103w   Homo sapiens gene for caspase-IO, exon I I and complete cds   Legionella fallonii macrophage infectivity potentiator (migh) gene,	
3954 M32666 Homo sapiens TTGB2 gene, intron 1, partial sequence 3955 Z72888 S.ccrevisiae chromosome VII reading frame ORF YGR103w 4B038979 Homo sapiens gene for caspase-10, exon 11 and complete eds Legionella fallonii macrophage infectivity potentiator (mip) gene,	0.056
3954 M32666 Homo sapiens TTGB2 gene, intron 1, partial sequence 3955 Z72888 S.ccrevisiae chromosome VII reading frame ORF YGR103w 4B038979 Homo sapiens gene for caspase-10, exon 11 and complete eds Legionella fallonii macrophage infectivity potentiator (mip) gene,	
3955 Z72888 S.ccrevisiae chromosome VII reading frame ORF YGR103w - 3956 AB038979 Homo sapiens gene for caspase-10, exon 11 and complete cds Legionella fallonii macrophage infectivity potentiator (mip) gene,	0.17
3956 AB038979 Homo sapiens gene for caspase-10, exon 11 and complete cds  Legionella fallonii macrophage infectivity potentiator (mip) gene,	2
Legionella fallonii macrophage infectivity potentiator (mip) gene,	0.009
	0.001
3958 AF148987 partial cds 3959 AF159173 Gallus gallus structural muscle protein titin mRNA, partial cds	0.075
3959 AF159173 Gallus gallus structural muscle protein titin mRNA, partial cds  Mycoplasma pulmonis FisZ (fisZ) gene, complete cds, methionyl-	1.2
3960 U34931 tRNA synthetase (metG) gene, partial cds	0.21
3961 AF072439 Rattus norvegicus zinc-finger protein-37 mRNA, complete cds	0.21
3962 AF079877 Mus musculus cyclin G2 (Ccng2) gene, complete cds	0.049
3963 X87947 S. cerevisiae ALG2 gene	0.049
Sectional Field Bone	0.024
Mus musculus, proteasome (prosome, macropain) subunit, beta type	
3964 BC004730 10, clone MGC:5837 IMAGE:3583052, mRNA, complete cds	6.8
3965 BC010610 Homo sapiens, clone IMAGE:4214515, mRNA, partial cds	2E-11
Mus musculus 10 days embryo cDNA, RIKEN full-length enriched	
3966 AK012050 library, clone:2610319A01, full insert sequence	
Homo sapiens, chromosome 4 open reading frame 1, clone	e-144
3967 BC000240 IMAGE:3352004, mRNA	
3969 AE007346 Streptococcus pneumoniae section 29 of 194 of the complete genome	e-144

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			1
NO	ACCESSN	DESCRIP	P VALUE
3970	AK021969	Homo sapiens cDNA FLJ11907 fis, clone HEMBB1000059	0.005
3971	U05765	Human annexin V (ANX5) gene, exon 7	4.9
3972	S68117	rPLP-A=prolactin-like protein A {5' region, exon 1, intron 1} [rats, Genomic, 1187 nt]	0.11
		Human DNA sequence from clone RP1-317N9 on chromosome 6,	
3973	AL358852	complete sequence [Homo sapiens]	0.18
3974	M26221	African green monkey origin of replication (ORS8) region	0.01
		Mouse MHC class I Lyt-2-a gene encoding lyt-2.1 T-cell surface	}
3975	M22064	alloantigen, complete cds	0.71
3976	AY039917	Arabidopsis thaliana putative wall-associated kinase 1 (F16F4.6) mRNA, complete cds	3.5
3977	U67488	Methanococcus januaschii section 30 of 150 of the complete genome	0.021
-		Homo sapiens N-acetylated alpha-linked acidic dipeptidase 2	
3979	XM_027089		1E-09
3980	AF062751	Manduca sexta soluble guanylyl cyclase beta-1 subunit mRNA, complete cds	0.53
1 1		Homo sapiens genomic DNA, chromosome 21q22.1, D21S226-AML	
3981	AP000214	region, clone f43D11-119B8, segment 12/12, complete sequence	0.0003
		Arabidopsis thaliana putative MLH1 protein (AT4g09140) mRNA,	
3982	AF360278	complete cds	0.0001
		2 Homo sapiens concentrative nucleoside transporter (CNT1) gene,	
3983	AF187968	exon 2	0.085
		Drosophila melanogaster H.M.S. Beagle transposon long terminal	
		repeat, complete sequence; and heat shock protein Hsp70Ab gene,	
3984	AY032741	promoter and partial cds	0.6
3985	NM_008697	Mus musculus ninein (Nin), mRNA	4
3986	AB055369	Macaca fascicularis brain cDNA, clone:QfIA-12661	0
3987	X15999	Kluyveromyces lactis mitochondrial tRNA-Val, COII and COI (partial)	0.023
3787	A13999	Homo sapiens chromosome 5 clone CTD-2178M23, complete	0.023
3988	AC008855	sequence	0.56
3989	X79482	I. punctatus Immunoglobulin mu heavy chain gene	0.00007
- 3707	2017402	Dalbulus charlesi NADH dehydrogenase subunit 1 gene.	0.00007
3990	AF051289	mitochondrial gene encoding mitochondrial protein, complete eds	0.072
3991	BC004382	Homo sapiens, clone IMAGE:3640982, mRNA, partial cds	e-147
		Homo sapiens hypothetical gene supported by AL117603	- V-1.
3992	XM 041432	(LOC91908), mRNA	1.3
		Mus musculus RIKEN cDNA 2310012P17 gene (2310012P17Rik),	
3993	NM_027142		0.027
		Oryza sativa rpl12-1 gene for chloroplast ribosomal protein L12,	
3994	AB022673	complete cds	0.69
		Bombyx mori nuclear polyhedrosis virus gene for putative DNA-	
3995	AB009987	directed RNA polymerase component lef8, complete cds	0.2
3996	NM_000144	Homo sapiens Friedreich ataxia (FRDA), mRNA	0.008

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#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE Homo sapiens LBP protein; likely ortholog of mouse CRTR-1 (LBP-3997 XM 002395 9), mRNA 0.58 Brassica napus cultivar Darmor AP31 gene, AP31-DN allele, partial 3998 AF056532 S.cerevisiae chromosome VII reading frame ORF YGR103w 3999 Z72888 0.008 Human DNA sequence from clone RP11-353M9 on chromosome 10. 4000 AL512660 complete sequence [Homo sapiens] 5.3 Cloning vector pVO205 hygromycin-B-phosphotransferase (hph) 4001 AF190131 gene, complete cds 1.7 4002 AJ132557 Oryctolagus cuniculus CYP19 gene, ovarian promoter region 0.069 AB001901 4003 Homo sapiens PACE4 gene, exon 4-7 4 4004 AE006784 Sulfolobus solfataricus section 143 of 272 of the complete genome 0.19 4005 XM_049237 Homo sapiens KIAA0841 protein (KIAA0841), mRNA 17 4006 AF107676 Aedes aegypti clone 416 Feilai family of SINES 0.54 4007 AB053857 TT virus gene for ORF1, partial cds, clone:Pd-10-3 0.24 Homo sapiens 3 BAC PAC-56F11 (Roswell Park Cancer Institute 4008 AC078798 Human BAC Library) complete semence 5 Mus musculus RIKEN cDNA 2810036K01 gene (2810036K01Rik), 4009 NM 026040 mRNA 2E-33 4011 XM_027300 Homo sapiens DKFZP434K114 protein (DKFZP434K114), mRNA 4E-08 Oryza sativa microsatellite MRG4298 containing (TA)X104, closest 4012 to marker S10620, genomic sequence AY021973 0.0009 Fruitfly strain g20 mitochondrial DNA, A+T-rich region, partial 4013 AB003097 sequence 0.000003 4014 AJ000682 Bos taurus PPARG gene, intron 1, 5' Helicobacter pylori, strain J99 section 65 of 132 of the complete 4015 AE001504 0.64 Homo sapiens hypothetical protein DKFZp761J17121 4016 NM_031442 (DKFZP761J17121), mRNA 0.007 Rattus norvegicus liver microsomal carboxylesterase mRNA, complete 4017 U10698 0.21 Homo sapiens putative chloride channel gene (CLCN6), exons 14, 15, 4018 AF009255 16, and 17 1.7 4019 NM 010657 Mus musculus kappa B and Rss recognition component (Krc), mRNA 0.17 Oryza sativa microsatellite MRG3425 containing (TA)X19, closest to 4020 AY021100 marker L246, genomic sequence 47 4021 U43092 Rattus norvegicus surfactant protein-A (SP-A) gene, complete cds 0.9 4022 XM_050127 Homo sapiens oligophrenin 1 (OPHN1), mRNA 0.64 Escherichia coli O157:H7 EDL933 genome, contig 3 of 3, section 195 4023 AE005576 of 290 0.45 4024 AF288693 Mus musculus Ubell (Ubell) gene, partial cds 0.0008 Taphozous sp. Brca1 (Brca1) gene, partial cds 4025 AF203748

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		Tuble 612 Treatest Heighbor (Diaster va. Genbank)	
NO	ACCESSN	DESCRIP	P VALUE
4026	AF179234	Homo sapiens short-chain dehydrogenase/reductase 1 (SDR1) gene, exon 1	0,56
4027	AF163735	Illicium parviflorum internal transcribed spacer 1, 5.8S ribosomal RNA gene, and internal transcribed spacer 2, complete sequence	0.62
4028	NM 026233	Mus musculus RIKEN cDNA 4933434I20 gene (4933434I20Rik), mRNA	6.5
		Human DNA sequence from clone CTA-191D12 on chromosome	6.5
		22q13.1 Contains two exons of the APOL2 gene for apolipoprotein L	
4029	AL031426	2, ESTs and GSSs, complete sequence [Homo sapiens]	0.000001
4030	XM 008148	Homo sapiens myeloid/lymphoid or mixed-lineage leukemia (trithorax (Drosophila) homolog); translocated to, 6 (MLLT6), mRNA	
4030	ANI_008146	Chaetopterus variopedatus histone H4 and histone H2A genes,	0.63
4031	AF007904	complete cds	0.009
1			
		Homo sapiens C-type (calcium dependent, carbohydrate-recognition	
4032	XM_051353	domain) lectin, superfamily member 5 (CLECSF5), mRNA	0.69
4033	U88154	Homo sapiens proline and glutamic acid rich nuclear protein isoform mRNA, partial cds	0.074
		Arabidopsis thaliana chromosome III P1 MJM20 genomic sequence,	0.077
4034	AC023838	complete sequence	0.2
4035	X15894	Sinapsis alba cab-1 gene for chlorophyll a/b-binding polypeptide	0.1
4036	X54742	Nicotiana plumbaginifolia beta-(1,3)-glucanase gene for a vacuolar isoform	1.8
4037	NM_017590	Homo sapiens hypothetical protein DKFZp434K0920 (DKFZp434K0920), mRNA	0.13
		Equus caballus interleukin-1 receptor antagonist (EqIL-1RA) mRNA.	0.13
4038	U92482	complete cds	0.38
4039	U47540	Aspergillus nidulans nitrogen regulatory protein (tamA) gene, complete cds	0.18
4040	AK001076	Homo sapiens cDNA FLJ10214 fis, clone HEMBA1006530	0.0000009
4041	AC002049	Homo sapiens Chromosome 22q11.2 Cosmid Clone 107d7 In BCRL2- GGT Region, complete sequence	1.2
	110002017	Dictyostelium discoideum glyceraldehyde-3-phosphate dehydrogenase	1.3
4042	U55243	(GAPDH) gene, partial cds	0.23
4043	Z16701	H. sapiens (D1S218) DNA segment containing (CA) repeat; clone AFM157xe7; single read	0,002
4044	AB012733	Rhododendron semibarbatum chloroplast matK gene for ribosomal maturase, complete cds	0.024
4045	AF034950	Bos taurus gonadotropin hormone receptor (GnRH) gene, partial eds	0.22
		Mus musculus mRNA for putative ubiquitin-specific protease (Usp9y	0.22
4046	AJ307017	gene)	1.7
4047	AE001159	Borrelia burgdorferi (section 45 of 70) of the complete genome	0.89
4048	AF239215	Forficula auricularia microsatellite FA2 sequence	0.0002

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Homo sapiens genomic DNA, chromosome 21q22.1, D21S226-AML   IE-18				
NO   ACCESSN   DESCRIP   P VALUE			Table 3A Nearest Neighbor (BlastN vs. Genbank)	
Homo sapiens genomic DNA, chromosome 21q22.1, D21S226-AML region, clone.B762015, complete sequence		ACCECON	DEG CO. III	
AP000249   region, clone:B762015, complete sequence	NO	ACCESSIV		P VALUE
Homo sapiens genomic DNA, chromosome 21q22.1, D21S226-AML   Fogion, clone:B762O15, complete sequence   D.61	4040	AD000240	Fromo sapiens genomic DNA, chromosome 21q22.1, D21S226-AML	
AF000249   region, clone B762015, complete sequence   AF35470   Mus musculus systetine dioxygenase gene, exon 2   Genetic and the sequence   AF35470   Mus musculus cystetine dioxygenase gene, exon 2   Genetic and the sequence   Genetic	4049	AF000249	region, cione:B/62O15, complete sequence	1E-18
AF355470	4050	A D0000240	Promo sapiens genomic DNA, chromosome 21q22.1, D21S226-AML	
Homo sapiens, hypothetical protein FLI13612 similar to COP9 (constitutive photomorphogenic), subunit 7b (Arabidopsis), clone MGC:16836 IMAGE:3893022, mRNA, complete cds library, clone-4993145012, full insert sequence   1.5			region, clone:B/62O15, complete sequence	
Constitutive photomorphogenic), subunit 7b (Arabidopsis), clone	4031	AF333470	Mus musculus cysteine dioxygenase gene, exon 2	0.61
MGC:16836 IMAGE:3893022, mRNA, complete cds			Homo sapiens, hypothetical protein FLJ12612 similar to COP9	
Must musculus adult male testis cDNA, RIKEN full-length enriched border   1.5	4052	PC010720	(Consultative photomorphogenic), subunit /b (Arabidopsis), clone	
AK016325   Library, clone:49932415G12, full linsert sequence	4032	BC010/39	MIGC.10836 INIAGE:3893022, mRNA, complete cds	4.5
April	4052	AT/016526	With musculus adult male testis cDNA, RIKEN full-length enriched	
4055				
Candidatus Carsonella ruddii natural-host Tainarys sordida RNA polymerase beta subunit (rpoB) and RNA polymerase beta subunit (rpoB) and RNA polymerase beta-prime subunit (rpoC) genes, partial ods   0.067				
AP268053   subunit (rpoB) and RNA polymerase beta-prime   subunit (rpoB) and RNA polymerase beta-prime   subunit (rpoC) genes, partial cds   0.067	4033	000038	Caeliornaudius elegans cosmid 121D11, complete sequence	4.3
AC016694			Candidatus Carsonella ruddu natural-host Tainarys sordida RNA	ĺ
AC016694   Hiomo sapiens BAC clone RP11-123G1 from Y, complete sequence   0.2	1056	A E1 < 90 62		
Mus musculus adult male testis cDNA, RIKEN full-length enriched   0.15	4030	A1 200033	subtlift (rpoc.) genes, partial cus	0.067
Mus musculus adult male testis cDNA, RIKEN full-length enriched   0.15	4057	AC016604	Homo coniens BAC clone DB11 122C1 6 V	
4058   AK019557   library, clone;4930402H05, full insert sequence   0,15   4059   AF23398   Escherichia coli strain KI1218 BfpA, (bfpA) gene, partial cds   0,663   4060   AK026269   Homo sapiens cDNA: FLJ22616 fis, clone HSi05164   5.7   4061   Z80362   Hemp sapiens cDNA: FLJ22616 fis, clone HSi05164   5.7   4062   XM 046282   (LOC92635), mRNA   0,007   4063   U83303   Local time 1- reverse transcriptase gene, partial cds, and granulocyte chemotactic protein-2 (GCP-2) gene, complete cds   4064   AF276169   Scaphirhynchus platorynchus microsatellite Spl-100 sequence   0,074   4065   AF156797   GAEP-alpha subunit gene, promoter and 5' UTR, and 4065   AF166797   GAEP-alpha subunit gene, promoter and 5' UTR, partial sequence   1.2   4066   AF110420   Mus musculus hoTgN37INRA locus sequence   4E-10   4067   AK021659   Homo sapiens cDNA FLJ11597 fis, clone HEMBA 1003856   6E-10   4068   AL512724   DKFZp547M202)   0,2   4069   S54531   L-plastin (exon 1, promoter) [human, Genomic, 3231 nt]   0,023   4070   L02110   Mus musculus and terminal retroviral insertion in the cCMP-phosphodiesterase (rd beta PDE) gene, intron 1, with the proviral insert encompassing the can yseudogene (3' end) and 3' LTR   4071   XM_015244   mRNA   Mus musculus adult male testis cDNA, RIKEN full-length enriched   4072   AK015420   library, clone:4930448118, full insert sequence   0,003	-1007	110010054	Mus musculus adult mala tartia DNA DESTRICTION 1, complete sequence	0.2
AF233898   Escherichia coli strain KII218 BfpA (bfpA) gene, partial cds   0.063	4058	AK010557	library clane:4020402105 6.11 incert as a suit length enriched	
AK026269   Homo sapiens cDNA: FLJ22616 fis, close HSi05164   5.7			Englarishia adi atraia KT1019 De. A. (C. A.)	
4061   Z80362			Homo coniene aDNA, EL 192616 E. alara ITO 5164	
Homo sapiens lrypothetical gene supported by AK026722   0.007			Li carriano LII. A. DDD manuda mana a la la la la la la la la la la la la	
Mode   March   7001	200302		1.7	
Human line-1 reverse transcriptase gene, partial cds, and granulocyte chemotactic protein-2 (GCP-2) gene, complete cds   5E-12	4062	YM 046282		
4063   U83303   chemotactic protein-2 (GCP-2) gene, complete cds   5E-12	1002	7111_040202		0.007
AF276169   Scaphiritynchus platorynchus microsatellite Spi-100 sequence   0.074	4063	TI83303	chemotoctic protein-2 (GCD-2) gene, complete ada	5T 10
Mus musculus ATP synthase coupling factor 6 gene, 5' UTR; and GABP-alpha subunit gene, promoter and 5' UTR, partial sequence   1.2			Scanhirhynchus platorynchus microcatellita Spi 100 common	
4065         AF156797         GABP-alpha subunit gene, promoter and 3' UTR, partial sequence         1.2           4066         AF110420         Mos musculus hoTgN3'INRA locus sequence         4E-10           4067         AK021659         Homo saplens cDNA FLJ11597 fis, clone HEMBA1003856         6E-10           4068         AL512724         Homo saplens mRNA; cDNA DKFZp547M202 (from clone         0.2           4069         S54531         L-plastin (exon 1, promoter) [human, Genomic, 3231 nt]         0.023           Mus musculus proviral retroviral insertion in the cGMP-phosphodiseterase (abe 2 PDE) gene, intro n 1, with the proviral insert encompassing the env pseudogene (3' end) and 3' LTR         2.1           4071         XM_015244         mRNA         1.5           4072         AK015420         Mis musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930448118, full insert sequence         0.003			bedynthylicatus pantorylicatus interosaterine Spi-100 sequence	0.074
4065         AF156797         GABP-alpha subunit gene, promoter and 3' UTR, partial sequence         1.2           4066         AF110420         Mos musculus hoTgN3'INRA locus sequence         4E-10           4067         AK021659         Homo saplens cDNA FLJ11597 fis, clone HEMBA1003856         6E-10           4068         AL512724         Homo saplens mRNA; cDNA DKFZp547M202 (from clone         0.2           4069         S54531         L-plastin (exon 1, promoter) [human, Genomic, 3231 nt]         0.023           Mus musculus proviral retroviral insertion in the cGMP-phosphodiseterase (abe 2 PDE) gene, intro n 1, with the proviral insert encompassing the env pseudogene (3' end) and 3' LTR         2.1           4071         XM_015244         mRNA         1.5           4072         AK015420         Mis musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930448118, full insert sequence         0.003			Mus musculus ATP synthese counting feater 6 game 51 IETP.	
4066	4065	AF156797	GABP-ainha subunit gene promoter and 5' LITP portial segments	
4067			Mus musculus hoToN37INRA locus seguence	
Homo supiens mRNA; cDNA DKFZp547M202 (from clone   0.2	4067		Homo sapiens cDNA FLI11597 fis. clone HEMBA 1003856	
4068         AL512724         DKFZp547M202)         0.2           4069         S54531         L-plastin (exon 1, promoter) [human, Genomic, 3231 nt]         0.023           Mus musculus proviral retroviral insertion in the cGMP-phosphediesterase (rd beta PDE) gene, intron 1, with the proviral insert encompassing the envy seudogene (3' end) and 3' LTR         2.1           4070         LO2110         Homo sapiens ring finger protein 21, interferon-responsive (RNF21), and mRNA         1.5           4071         XM_015244         mRNA         1.5           4072         AK015420         library, clone:4930448118, full insert sequence         0.003			Homo saniens mRNA: cDNA DKEZp547M202 (from elene	0E-10
4069   S34531   L-plastin (exon 1, promoter) [human, Genomic, 3231 nt]   0.023	4068	AL512724	DKFZp547M202)	0.2
Mus musculus proviral retroviral insertion in the cGMP- phosphodiesterase (rd beta PDE) gene, intron 1, with the proviral insert encompassing the eavy pseudogene (3° end) and 3′ LTR 2.1   Homo sapiens ring finger protein 21, interferon-responsive (RNF21),   mRNA   Mus musculus adult male testis cDNA, RIKEN full-length enriched     dozen	4069		L-plastin (exon 1, promoter) [human Genomic 3231 nt]	
4070   L02110   phosphodiesterase (rd beta PDE) gene, intron 1, with the proviral insert encompassing the eary pseudogene (3' end) and 3' LTR   2.1			Mus musculus provinal retroviral insertion in the cGMP.	0.023
4070         L/02110         insert encompassing the eary pseudogene (3° end) and 3° LTR         2.1           4071         XM_015244         Homo sapiens ring finger protein 21, interferon-responsive (RNF21), mRNA         1.5           4072         AK015420         Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930448118, full insert sequence         0.003	- 1		phosphodiesterase (rd beta PDE) gene intron 1 with the provint	1
Homo sapiens ring finger protein 21, interferon-responsive (RNF21),   Homo sapiens ring finger protein 21, interferon-responsive (RNF21),   1.5   Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930448118, full insert sequence   0.003	4070	L02110	insert encompassing the environment (3' end) and 3' I TP	21
4071 XM_015244 mRNA 1.5  Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone;4930448118, full insert sequence 0.003			Homo sapiens ring finger protein 21 interferon-responsive (PNE21)	2.1
Mis musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930448118, full insert sequence 0.003	4071	XM 015244	mRNA	15
4072 AK015420 library, clone:4930448118, full insert sequence 0.003			Mus musculus adult male testis cDNA RIKEN full-length enriched	
	4072	AK015420	library, clone:4930448118, full insert sequence	0.003
and I common has				3.005
4073 U67583 Methanococcus jannaschii section 125 of 150 of the complete genome 0.021	4073	U67583	Methanococcus jannaschii section 125 of 150 of the complete genome	0.021
5.551			Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar Salar	
4074 X05640 Mouse NF-M gene for middle-molecular-mass neurofilament protein 0.072	4074	X05640	Mouse NF-M gene for middle-molecular-mass neurofilament protein	0.072

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#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEQ ID NO ACCESSN DESCRIP P VALUE Anthocidaris crassispina mRNA for dynein beta-heavy chain, 4075 D01021 complete cds 0.61 4076 XM 029072 Homo sapiens LOC89256 (LOC89256), mRNA 0.006 4077 NM 017757 Homo sapiens hypothetical protein FLJ20307 (FLJ20307), mRNA 0.26 Nicotiana tabacum calcium/calmodulin-dependent protein kinase 4078 AF145592 (CCaMK) gene, exons 0.002 Arabidopsis thaliana somatic embryogenesis receptor-like kinase 3 4079 AF384970 (SERK3) mRNA, complete cds 5.3 Dictyostelium discoideum developmental protein DG1037 (DG1037) 4080 AF238313 gene, partial cds 0.2 Rattus norvegicus Ca++/calmodulin-dependent protein kinase II. delta 4081 NM 012519 subunit (Camk2d), mRNA 0.071 Human DNA sequence from clone RP11-349A16 on chromosome 4082 AL590384 Xq22.3-24, complete sequence [Homo sapiens] 0.11 4083 AF391284 Homo sapiens 11p15.5 clone LOH11A, partial sequence 0.17 4086 XM 034197 Homo sapiens KIAA1505 protein (KIAA1505), mRNA e-153 Sinorhizobium meliloti plasmid pSymA section 61 of 121 of the 4087 AE007255 complete plasmid sequence 5.4 Streptococcus pyogenes M1 GAS strain SF370, section 115 of 167 of 4088 AE006586 the complete genome 16 4089 AF391284 Homo sapiens 11p15.5 clone LOH11A, partial sequence 0.14 U41549 4090 Caenorhabditis elegans cosmid F22F1 0.065 Pseudosymblepharis schimperiana tRNA-Leu, partial sequence: trnLtrnF intergenic spacer, complete sequence; and tRNA-Phe, partial 4091 AF231164 sequence; chloroplast genes for chloroplast products 0.066 4092 XM 005131 Homo sapiens chromatin accessibility complex 1 (CHRAC1), mRNA 0.39 Mus musculus RIKEN cDNA 2010109I03 gene (2010109I03Rik). 4093 NM 025929 mRNA 5.1 4094 D.discoideum mRNA for calcium binding protein X82784 0.00001 Caedibacter taeniospiralis 47 R body synthesis and assembly (rebA rebB, rebC, rebD) genes, complete cds 4095 U04524 0.58 XM 050063 4096 Homo sapiens B-cell CLL/lymphoma 9 (BCL9), mRNA 4097 XM 037650 Homo sapiens syndecan 1 (SDC1), mRNA 2 Homo sapiens retinitis pigmentosa GTPase regulator (RPGR) gene, 4098 AF286472 exon ORF15 partial cds 0.076 4099 L05466 Pneumocystis carinii beta-tubulin gene, complete cds 0.009 Human DNA sequence from cosmid U238E5, between markers 4100 Z69723 DXS6791 and DXS8038 on chromosome X 0.18 Human DNA for CAAF1 (calcium-binding protein in amniotic fluid 4101 D83657 1), complete cds 5E-30 Botrytis cinerea strain T4 cDNA library under conditions of nitrogen 4102 AL117050 deprivation 2.1 Vibrio cholerae chromosome I, section 76 of 251 of the complete 4103 AE004168 chromosome 2.3

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#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE 4104 BC007114 Homo sapiens, clone IMAGE:4295422, mRNA Penicillium chrysogenum mitochondrion genes for rRNA, tRNA and 4105 D13859 S5 ribosomal protein, complete cds 0.75 4106 AF137068 Canis familiaris cubilin (CUBN) mRNA, complete cds 4.3 Mus musculus 11 days embryo cDNA, RIKEN full-length enriched 4107 AK012387 library, clone:2700047H13, full insert sequence 0.38 NC 001942 4108 Mycoplasma arthritidis bacteriophage MAV1, complete genome 4109 AF333431 Homo sapiens pseudo attP site psiD, partial sequence 0.45 4110 AF305628 Cubitermes subarquatus microsatellite P19 sequence 0.68 C.ellipsoidea rbcL, rps14, trnM, trnG, trnD, trnS, rps4, atpE, atpB 4111 D10997 genes, complete cds 1.8 Human DNA sequence from clone LL22NC01-75B8 on chromosome 4112 Z99704 22 Contains GSSs, complete sequence [Homo sapiens] 0.005 Homo sapiens androgen-induced prostate proliferative shutoff 4113 XM 016579 associated protein (KIAA0979), mRNA 1.6 Mus musculus glycerol-3-phosphate acyltransferase, mitochondrial NM 008149 (Gpam), mRNA 55 4115 XM 036116 Homo sapiens KIAA0594 protein (KIAA0594), mRNA e-144 Homo sapiens microfibrillar-associated protein 2 (MFAP2), mRNA 4116 XM 030510 2.4 4117 AB010203 Leptospira interrogans gene, 19kb region containing 5S rRNA gene 1.4 4118 XM_007094 Homo sapiens collagen, type IV, alpha 1 (COL4A1), mRNA Fugu rubripes double stranded RNA adenosine deaminase RED1B 4120 AF124049 gene, complete cds 0.066 4121 AY007236 Pneumocystis carinii pheromone receptor a mRNA, complete cds 0.53 Aptostichus simus clone PD1 16S ribosomal RNA gene, partial 4122 AF307966 sequence: mitochondrial gene for mitochondrial product 2 4123 AJ001022 Marmata monax mRNA for tumor suppressor, p53 1.3 Polycycnis gratiosa maturase (matK) gene, partial cds; chloroplast 4124 AF239469 gene for chloroplast product 0.65 4125 X75675 C.parapsilosis mitochondrial ND6 and ND1 genes 0.00001 Glycine max chitinase class I (Chia1) and hypothetical protein genes, 4126 AF335589 complete cds 0.0003 4127 U33175 Oryza sativa sucrose phosphate synthase gene, complete cds 6.4 4128 AB028198 Arabidopsis thaliana ZF14 mRNA, complete cds 0.008 4129 AF372653 Heteropneustes fossilis prolactin hormone (PRL) mRNA, complete cds 1.3 4130 M63955 S.oedipus MHC class I SO-NI gene 0.6 Mus musculus, protein kinase, cAMP dependent, catalytic, alpha, 4131 BC003238 clone MGC:6169 IMAGE:3497908, mRNA, complete cds 0.22 AJ001688 Homo sapiens NKG2D gene, exons 6-9 0.67 4133 D10872 Humam h NAT allele 3-2 gene for arylamine N-acetyltransferase 0.016

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			T
NO	ACCESSN	DESCRIP	P VALUE
		Cotesia congregata EP1 locus with complete integrated Cotesia	
4134	AF049877	congregata virus EP1 circle and partial integrated A circle	0.19
4135	Y15783	Rickettsia prowazekii groESL gene	0.3
4136	Y00354	Xenopus laevis gene encoding vitellogenin A2	0.077
4137	NC_001950	Scyliorhinus canicula mitochondrion, complete genome	0.019
		Macromia splendens small subunit ribosomal RNA gene, partial	1
4138	AF266048	sequence; mitochondrial gene for mitochondrial product	0.42
		Human DNA sequence from clone 370N13 on chromosome Xq25-	
		26.3. Contains an exon of the GRIA3 gene for glutamate receptor,	l
4139	AT 025426	ionotrophic, AMPA 3. Contains ESTs, complete sequence [Homo	
4139	AL035426	sapiens]	0.0006
4140	AF018434	Mus musculus long-chain acyl-CoA dehydrogenase (Acadl) gene, intron 1	
4140.	AF010434	Mus musculus U2AF small subunit-related protein (U2af1-rs1) gene,	6.4
4141	AF309654	complete cds	0.64
4142	AJ009034	Bos taurus plp gene	2.1
4143	XM 028213	Homo sapiens up-regulated by BCG-CWS (LOC64116), mRNA	0.078
1113	7LIVI_020213	Tionio sapiciis up-regulated by Beo-ews (EOco+116), ilikava	0.078
4144	U67550	Methanococcus jannaschii section 92 of 150 of the complete genome	0.081
4145	AB037771	Homo sapiens mRNA for KIAA1350 protein, partial cds	0.081
12.15	120031111	Homo sapiens genomic DNA, chromosome 21q22.2, clone:T695,	<del></del>
4146	AP001438	LB7T-ERG region, complete sequence	0.009
		Callithrix geoffroyi clone CGE85 mitochondrial D-loop, partial	0.007
4147	U88995	sequence	0.57
4148	AE007343	Streptococcus pneumoniae section 26 of 194 of the complete genome	1.9
4149	Z83832	Avena sativa mRNA for UDP-glucose:sterol glucosyltransferase	0.072
		Homo sapiens, hepatocellular carcinoma-associated antigen 59, clone	
4150	BC007664	MGC:866 IMAGE:3140172, mRNA, complete cds	0
4151	NM_002577	Homo sapiens p21 (CDKN1A)-activated kinase 2 (PAK2), mRNA	5E-39
		Homo sapiens ADP-ribosylation factor GTPase activating protein 1	
4152	XM_040120		2.1
		Rattus norvegicus Insulin-like growth factor II (somatomedin A)	
4153	NM_031511	(Igf2), mRNA	0.024
4154	XM_049817	Homo sapiens Sec23-interacting protein p125 (P125), mRNA	1.8
		Homo sapiens, clone MGC:13350 IMAGE:4333615, mRNA,	
4155	BC010892	complete cds	0.000004
1100	A T000707	Hordeum vulgare mRNA for alpha-keto acid dehydrogenase-like	
4156	AJ222787	protein, clone RG136	0.075
		W Davi	
41.57	AT 021701	Human DNA sequence from clone CTB-1109B5 on chromosome 22	0.00000:
4157	AL021306	Contains a GSS, complete sequence [Homo sapiens]	0.000001
4158	AF315813	Norwalk virus (Hu/NLV/OC96065/1996/JP) polymerase gene, partial	
4138	AF315813	cds	0.71

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
4159	AE001407	Plasmodium falciparum chromosome 2, section 44 of 73 of the complete sequence	0.0008
4160	U32706	Haemophilus influenzae Rd section 21 of 163 of the complete genome	0.6
4161	AF293663	Anopheles culicifacies 12S ribosomal RNA gene, partial sequence, mitochondrial control region, complete sequence; and tRNA-Ile gene,	
4101	AF293003	partial sequence; mitochondrial genes for mitochondrial products	0.066
4162	XM_037617	Homo sapiens acetyl-Cocnzyme A carboxylase alpha (ACACA), mRNA	3E-35
4163	AF305712	Bos taurus insulin-like growth factor binding protein-3 (IGFBP3) gene, complete cds	0.23
		Mus musculus adult male cecum cDNA, RIKEN full-length enriched	
4164	AK018618	library, clone:9130014E20, full insert sequence	0.68
4165	Y00354	Xenopus laevis gene encoding vitellogenin A2	0.079
4166	M61827	Human leukosialin (CD43) gene, complete cds	2.2
4167	M64861	D.discoideum TFIID mRNA, complete cds	0.066
4169	AB028894	Mus musculus Rps11, U35 genes for ribosomal protein S11 and U35 snoRNA, complete cds and sequence	0.6
4170	XM_003974	Homo sapiens hypothetical protein FLJ11159 (FLJ11159), mRNA	0.68
4171	AF100658	Caenorhabditis elegans cosmid H08G01	0.75
		Sedum multiceps maturase K (matK) gene, partial cds; chloroplast	
4172	AF115645	gene for chloroplast product	0.0001
4173	AF064029	Helianthus tuberosus lectin 1 mRNA, complete cds	0.11
4175	AF093754	Arabidopsis thaliana trans-membrane nitrate transporter protein AtNRT2:1 mRNA, complete cds	0.22
		Human L1Heg repetitive element from the intergenic region of the	
4176	M14292	epsilon and G-gamma globin genes	2E-08
4177	NM_013564	Mus musculus insulin-like 3 (Insl3), mRNA	0,067
4178	AF218815	Coturnix japonica early response to neural induction ERNI (ERNI) mRNA, partial cds	0.21
4179	U67616	Human 28S ribosomal RNA pseudogenes and alu repeat region sequence	6E-20
12/2		Frankia sp. Ari3 cryptic plasmid pFO31 ORF AF, korSAF gene, ORF	0L-20
1 1		CrF, ORF DrF, repF gene, ORF FrF, ORF GF, ORF HF, ORF JF,	
		ORF KrF, ORF LrF, ORF MF, ORF NrF, ORF OF, ORF PrF, parAF	
4180	AJ297945	gene and ORF SrF	0.68
4181	AB035429	Homo sapiens AGL gene for glycogen-debranching enzyme, exon 14	0.62
4182	AE006774	Sulfolobus solfataricus section 133 of 272 of the complete genome	0.063_
		Dictyostelium discoideum vacuolar proton ATPase 100-kDa subunit	
4183	U38803	(vatM) mRNA, complete cds	0.025
		Human macrophage colony stimulating factor receptor (c-fms) gene,	
4184	U78096	exon 1A, 2 and partial cds	0.0001

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#### Table 3A. Nearest Neighbor (BlastN vs. Genbank) SEQ ID NO ACCESSN DESCRIP P VALUE 4185 X57037 T.thermophila G8-scRNA DNA 5.7 4186 AJ001005 Picea abies chloroplast DNA for rbcL pseudogene 0.67 Homo sapiens Apobec-1 complementation factor; APOBEC-1 4187 XM 048652 stimulating protein (ACF), mRNA 4E-12 4188 Y07654 P.crispum pal1 gene 0.24 4189 Z78708 H.sapiens flow-sorted chromosome 6 HindIII fragment, SC6pA14H12 1E-28 4190 AE002143 Ureaplasma urealyticum section 44 of 59 of the complete genome 2.1 4191 AF247729 Oncorhynchus mykiss glucose transporter 1B mRNA, partial cds 1.8 4192 Z72929 S.cerevisiae chromosome VII reading frame ORF YGR144w 0.003 4193 M87754 Human simple repeat polymorphism 0.081 4194 AL021809 S nombe chromosome I cosmid c25A8 4195 D12820 Rat mRNA for GUST27 protein, complete cds 37 4196 M60352 Mouse testosterone 16a-hydroxylase type b (16aoh-b), exon 1 4.7 4197 XM 007360 Homo sapiens hepatocyte nuclear factor 3, alpha (HNF3A), mRNA 0.95 4198 Z36064 S.cerevisiae chromosome II reading frame ORF YBR195c 0.52 4199 AK023959 Homo sapiens cDNA FLJ13897 fis, clone THYRO1001706 0.49 Dictyostelium discoideum rtoA gene, complete cds 4200 U48298 0.68 4201 L47461 Heterodontus francisci ter beta gene 16 4202 AB041023 Humulus lupulus DNA, 18S-26S intergenic spacer, complete sequence 0.0008 Hordeum vulgare high pI alpha-glucosidase (AGL97) gene, complete 4203 AF118226 1.8 Platybrachys decemmacula 16S ribosomal RNA gene, partial 4204 AF158049 sequence; mitochondrial gene for mitochondrial product 0.077 Drosophila virilis decapentaplegic protein (dpp) gene, complete cds, 4205 U63855 haploinsufficient region 0.2 Maoricicada cassiope isolate TB-MI-004 tRNA-Asp gene, complete sequence; ATPase subunit 8 gene, complete cds; and ATPase subunit 4206 AF248814 6 gene, partial cds; mitochondrial genes for mitochondrial products 0.023 4207 XM 037529 Homo sapiens KIAA0410 gene product (KIAA0410), mRNA 0.78 4208 AF330636 Plant DNA excision vector pX6-GFP, complete sequence 0.76 Homo sapiens mRNA; cDNA DKFZp586E1621 (from clone 4209 AL080235 DKFZp586E1621) 0.007 4210 NM 021262 Rattus norvegicus Acid phosphatase 1, soluble (Acpl), mRNA 0.16 4211 M11449 Yeast mitochondrial ori2-ori7 region DNA with putative peptide 0.25 Carsonella ruddii natural-host Ctenarytaina longicauda 16S ribosomal RNA gene, partial sequence; 23S ribosomal RNA and 5S ribosomal RNA genes, complete sequence: transaldolase (tal) and alkyl-4212 AF211134 hydroperoxide reductase small subunit (ahpC) genes, complete> 0.074 Sporodiniella umbellata translation elongation factor 1-alpha (EF-4213 AF157293 lalpha) gene, partial cds 0.77

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#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE Human DNA sequence from clone RPI-68PI5 on chromosome 11p13-14.2 Contains GSSs and ESTs. Contains part of a novel gene. complete sequence [Homo sapiens] 4214 AL390767 0.004 4215 XM 004579 Homo sapiens HGC6.1.1 protein (HGC6.1.1), mRNA 4E-08 Homo sapiens similar to KIAA0220 protein (H. sapiens) (LOC92263), 4216 XM 043964 5E-40 Human DNA sequence from clone RP1-68P15 on chromosome 11p13-14.2 Contains GSSs and ESTs. Contains part of a novel gene, 4217 AL390767 complete sequence [Homo sapiens] 0.004 4218 AF147082 Homo sapiens gamma-glutamyl hydrolase gene, exons 3 through 7 0.088 Oryza sativa microsatellite MRG5885 containing (TTA)X8, closest to 4219 AY023560 marker R2561, genomic sequence 0.026 Homo sapiens hepatocellular carcinoma-associated antigen 127 4220 XM 050088 (HCA127), mRNA 0.007 Plasmodium falciparum putative erythrocyte binding protein EBL-1 4221 AF131999 (ebl-1) gene, partial cds 0.68 Solanum fendleri farnesyl-protein transferase beta subunit gene. AF008434 4222 intron 10 0.21 4223 AF367267 Arabidopsis thaliana AT3g26020/MPE11 17 mRNA, complete cds 0.53 4224 XM_006706 Homo sapiens primase, polypeptide 1 (49kD) (PRIM1), mRNA 7E-12 Mus musculus adult male lung cDNA, RIKEN full-length enriched library, clone:1200009I20, full insert sequence 4225 AK004675 0.017 Arabidopsis thaliana genomic DNA, chromosome 5, BAC 4226 AB026660 clone:T20O7 0.23 4227 AF272387 Homo sapiens leucine zipper nuclear factor (BLZF1) gene, exon 6 5E-09 Mycoplasma gallisepticum (strain A5969) 16S-, 23S-, 5S ribosomal 4228 I 08897 RNA (rrsA, rrlA, rrfA) genes 0.073 4229 Z26314 P.falciparum gene for STARP antigen 0.008 4230 NC 001942 Mycoplasma arthritidis bacteriophage MAV1, complete genome 0.18 4231 Y18277 Gallus gallus mRNA for neurobeachin, partial 3 Homo sapiens pleiomorphic adenoma gene-like 2 (PLAGL2), mRNA 4232 XM 047009 1.8 4233 X84729 S.cereale repeated DNA sequence Human DNA sequence from clone RP11-460C6 on chromosome 9, 4234 AL513132 complete sequence [Homo sapiens] 0 E.herklotsi mitochondrial genes for NADH Dehydroenase subunit 5 4235 Z71694 0.6 4236 M.capricolum DNA for CONTIG MC294 0.54 Human DNA sequence from intron 22 of the factor VIII gene, Xq28. Contains the end of a 9.5kb repeated region, int22h-1, involved in 4237 X86012 many cases of haemophilia 0.0000005 Praon barbatum 16S ribosomal RNA gene, partial sequence: 4238 AF174355 mitochondrial gene for mitochondrial product 0.026

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
4239	AY039607	Arabidopsis thaliana AT3g52380/F22O6_240 mRNA, complete cds	6.1
4240	AJ223385	Fowlpox virus strain HP-440 DNA, isolate FP9, 14.6 kb fragment	0.000004
4241	AF277282	Daphnia cristata country Lithuania 12S ribosomal RNA gene, partial sequence	1.9
4242	NM_023850	Mus musculus carbohydrate (kcratan sulfate Gal-6) sulfotransferase 1 (Chst1), mRNA	0.59
4243	AJ238599	Lepilemur septentrionalis genomic fragment, RAPD with primer operon OPH4, clone A1	0.2
4244	XM_051747	Homo sapiens eukaryotic translation initiation factor 4 gamma, 2 (EIF4G2), mRNA	3E-58
4245	M63376	Human TRPM-2 protein gene, exons 1,2 and 3	0.7
4246	AF367267	Arabidopsis thaliana AT3g26020/MPE11_17 mRNA, complete cds	0.59
4247	AJ294714	Pichia anomala leu2 gene for beta-isopropylmalate dehydrogenase	0.17
4248	AF320616	Mus musculus amnionless precursor protein (Amn) gene, partial eds	0.57
4249	AL049609	S.pombe chromosome III cosmid c297	1.8
4250	AJ252313	Homo sapiens genomic hybrid Rhesus box	3.8
4251	XM_034693	Homo sapiens ectodermal dysplasia 1, anhidrotic (ED1), mRNA	0.2
4252	NC_001463	Caprine arthritis-encephalitis virus, complete genome	1,3
4253	AC018655	Homo sapiens 12q BAC RP11-946G22 (Roswell Park Cancer Institute Human BAC Library) complete sequence	5.9
4254	AF003519	Paraligoneurus sp. 16S ribosomal RNA gene, partial sequence	0.024
4255	AL049434	Homo sapiens mRNA; cDNA DKFZp586M151 (from clone DKFZp586M151)	0.0008
4256	U39321	Triticum aestivum acetyl-CoA carboxylase gene, exons 1-30, complete cds	0.22
4257	AB055106	Oryza sativa V-ATPase B gene for vacuolar ATPase B subunit, complete cds	1.7
4258	AF150091	Rattus norvegicus small zinc finger-like protein (TIM10) mRNA, complete cds	0.24
4259	AF047677	Homo sapiens dystrophin (DMD) gene, deletion breakpoints 1-3 in intron 5	0.025
4260	Z81504	Caenorhabditis elegans cosmid F15H9, complete sequence	0.24
4261	AF226147	HIV-1 isolate 630469c from USA protease (pol) gene, partial cds	2
4262	AF270072	Staphylococcus epidermidis strain SR1 clone step. 1047b05 genomic sequence	0.68
4263	U56088	Human periodic tryptophan protein 2 (PWP2) gene, exons 3 to 14	0.076
4264	X83692	Brassica napus (tournefortii) mitochondrial genes for atp6, and ORF263	2.1

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CEO TO		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
4265	AF243425	Penicillium oxalicum vacuolar serine protease mRNA, complete cds	0.63
4266	NM_024381	Rattus norvegicus ATP-stimulated glucocorticoid-receptor translocaton promoter (Gyk), mRNA	0.68
4267	AF098919	Gallus gallus alpha-globin gene domain 5' region	2.3
4268	AJ297909	Hypocrea jecorina rho3 gene, exons 1-5	0.23
4269	U67528	Methanococcus jannaschii section 70 of 150 of the complete genome	0.029
4270	AF104920	Arabidopsis thaliana BAC T9E19	0.026
4271	AF137394	Euperipatoides rowelli elongation factor-1 alpha mRNA, partial cds	0.23
		Homo sapiens intrinsic factor-vitamin B12 receptor (CUBN) gene,	
4272	AF243087	exons 11, 12, and 13	0
4273	AF264703	Astyanax mexicanus retinal homeobox protein (Rx1) mRNA, complete cds	2.2
		Chironomus pallidivittatus 4L ORF for putative recombinase	2.2
4274	AJ311054	subtelocentric clone, Cp5.5	0.025
		Candidatus Carsonella ruddii natural-host Calophya schini ATP	
		synthase alpha subunit (atpA) gene, partial cds; ATP synthase gamma	
		subunit (atpG) gene, complete cds; and ATP synthase beta subunit	
4275	AF267209	(atpD) gene, partial cds	0.0008
4276	Z17148	H. sapiens (D11S935) DNA segment containing (CA) repeat; clone AFM254zb9; single read	0.008
4270	Z1/140	Gallus gallus T cell receptor delta chain (TCRD) pseudogene, partial	0,008
4277	AF175434	sequence	2.3
		Mus musculus 10 days embryo cDNA, RIKEN full-length enriched	
4278	AK011996	library, clone:2610306H15, full insert sequence	0.49
		Staphylococcus epidermidis strain SR1 clone step.1005h04 genomic	
4279	AF269490	sequence	0.087
4280	XM_050118	Homo sapiens hephaestin (HEPH), mRNA	2E-75
4281	AF232828	Mus musculus ventral neuron-specific protein 1 NOVA1 (Nova1) mRNA, partial cds	0.000
4201	A('232020	Homo sapiens hypothetical gene supported by AK023162	0.008
4282	XM 029168	(LOC90120), mRNA	0.53
4283	L31645	Helianthus annuus ribosomal protein S3a mRNA, complete cds	0.67
4284	XM_012756	Homo sapiens transcription factor 4 (TCF4), mRNA	0.22
		Mus musculus adult male corpus striatum cDNA, RIKEN full-length	
4285	AK021049	enriched library, clone: C030006F08, full insert sequence	0.006
4286	AB020763	Ptychodera flava mRNA for Pax19, complete cds	0.26
4287	XM_047401	Homo sapiens G protein-coupled receptor 51 (GPR51), mRNA	0.25
4288	ATMENNER	Osmia satoi 16S ribosomal RNA gene, partial sequence;	
4288	AF250952 AF267985	mitochondrial gene for mitochondrial product	0.24
4289	AF267985 AJ409503	Homo sapiens 15 kDa selenoprotein gene, exon 4  Mus musculus RNA binding site for Dazl protein, clone jd8	0.024
7470			0.029

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	,	Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		Table 51k Treatest (Cignibol (Blastiv Vs. Gellbank)	
NO	ACCESSN	DESCRIP	P VALUE
4292	XM_050269		0.21
4293	AK004286	Mus musculus 18 days embryo cDNA, RIKEN full-length enriched library, clone:1110057K12, full insert sequence	2.1
4294	NC 002095	Chlorobium limicola plasmid pCL1, complete sequence	0.24
4295	AC006678	Caenorhabditis elegans cosmid R05G9, complete sequence	2.1
4296	AF217235	Staphylococcus aureus pathogenicity island SaPIbov, complete sequence	0.009
4297	XM_012045	Homo sapiens hypothetical protein FLJ13074 (FLJ13074), mRNA	0.68
4298	NM_022087		e-158
4299	XM_010678	Homo sapiens similar to zinc finger protein 258 (H. sapiens) (LOC65323), mRNA	7E-63
4300	AJ311847	Nicotiana tabacum chloroplast mRNA for FtsZ-like protein (ftsZ gene), clone FtsZ2-2	0.55
4301	Z24366	H. sapiens (D18S480) DNA segment containing (CA) repeat; clone AFM320yc9; single read	8E-12
4302	AK003805	Mus musculus 18 days embryo cDNA, RIKEN full-length enriched library, clone:1110019B22, full insert sequence	0.009
4303	AB047606	Macaca fascicularis brain cDNA, clone:QnpA-10881	0
4304	AK000493	Homo sapiens cDNA FLJ20486 fis, clone KAT08039	0.24
4305	AF375597	Mus musculus medium and short chain L-3-hydroxyacyl-Coenzyme A dehydrogenase (Mschad) gene, exons 2 through 8, and complete cds; nuclear gene for mitochondrial product	0.75
4306	AP000301	Homo sapiens genomic DNA, chromosome 21q22.1, D21S226-AML region, clone:D16B8, complete sequence	0.081
4307	AK019632	Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930455M05, full insert sequence	0.068
4308	X94556	R.cingulata DNA for internal transcribed spacer ITS2 of rDNA unit	0.079
4310	AY027174	Arabidopsis thaliana sequence flanking 3' end of Ds-GeneTrap insertion from line GT2850	0.2
4311	AC092063	Homo sapiens clone RP11-745H7, complete sequence	0.14
4312	AJ297034	Homo sapiens SDS-stable vimentin-bound DNA fragment HEF19VIM18	e-117
4313	Z82088	Caenorhabditis elegans cosmid ZK256, complete sequence	_0.015
4314	AC006709	Caenorhabditis elegans cosmid Y119C1A, complete sequence	0.17
4315	AF016620	Mus musculus Rb-8 neural cell adhesion molecule long form (RNCAM) mRNA, partial cds	6.2
		Mus musculus 10, 11 days embryo cDNA, RIKEN full-length	
4316	AK013121	enriched library, clone:2810421E14, full insert sequence	1.9
4317	XM_033379	Homo sapiens KIAA1607 protein (KIAA1607), mRNA	4E-09
4318	AL355864	Human DNA sequence from clone RP4-803A2 on chromosome 1, complete sequence [Homo sapiens]	1.4

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEO ID	<del>                                     </del>	Troublest Truginos (Diastry vs. Octoback)	
NO	ACCESSN	DESCRIP	P VALUE
		Plasmodium falciparum chromosome 2, section 59 of 73 of the	
4319	AE001422	complete sequence	0.028
	1	Homo sapiens mRNA; cDNA DKFZp434E1512 (from clone	
4320	AL157437	DKFZp434E1512); partial cds	4.9
4321	U31929	Human orphan nuclear receptor (DAX1) gene, complete cds	5.4
		Hylobates moloch NAN03 cytochrome b gene, partial cds, and tRNA-	
		Thr, tRNA-Pro, and tRNA-Phe genes, complete sequence;	l
4322	AF338875	mitochondrial genes for mitochondrial products	0.6
4323	X04502	Human SLPI gene for secretory leukocyte protease inhibitor	2
4324	U07018	Human immunodeficiency virus type 1 SH803 (env) gene, partial cds	0.64
4325	AF046143	Homo sapiens chromosome Xp22 958-13	0.023
4326	U17081	Human fatty acid binding protein (FABP3) gene, complete cds	6
1520	017001	Plasmodium falciparum partial mRNA for delta-aminolevulinic acid	
4327	AJ249365	dehydratase	0.009
4328	AF090936	Homo sapiens clone HQ0582	0.006
4329	X97970	A.thaliana mRNA for RNA helicase	0.079
4330	XM 047794	Homo sapiens KIAA1641 protein (KIAA1641), mRNA	e-156
4331	AF334172	Zca mays clone B7 chromosome 4 centromeric region	3.9
		Human immunodeficiency virus type 1 proviral partial gp160 gene for	
4332	AJ401041	envelope protein, strain J_97DC.KTB147	0.64
4333	XM_038175	Homo sapiens KIAA0872 protein (KIAA0872), mRNA	0.65
		Chlamydomonas reinhardtii chloroplast DNA for rps18 gene, and	
4334	Y16473	orf570	0.028
		Eucalyptus tetragona photosystem Q(B) protein (psbA) pseudogene	
		and tRNA-His (trnH) gene, partial sequence; chloroplast genes for	
4335	AF190381	chloroplast products	0.008
4224	1 Too (0.0)	Lactococcus lactis subsp. lactis IL1403 section 38 of 218 of the	
4336	AE006276	complete genome	0.22
4337	AK020646	Mus musculus adult male urinary bladder cDNA, RIKEN full-length	
4337	AK020040	enriched library, clone:9530077A04, full insert sequence	2.3
- 1		Human DNA sequence from cosmid 24F8 from a contig from the tip of the short arm of chromosome 16, spanning 2Mb of 16p13.3.	
4338	Z69666	Contains ESTs, repeat polymorphism and CpG island	0.010
1036	207000	contains 2018, repeat polymorphism and cpc island	0.018
4339	AF017254	West Nile virus nonstructural protein NS5 (NS5) gene, partial cds	0.73
4341	NM 008645	Mus musculus murinoglobulin 1 (Mug1), mRNA	0.00009
		0 ( 0,0)	
		Saccharomyces pastorianus CBS1538 small subunit ribosomal RNA	
4342	AF114927	gene, mitochondrial gene for mitochondrial RNA, complete sequence	0.25
		Human DNA sequence from clone CTD-2010K20 on chromosome 10,	
4343	AL451057	complete sequence [Homo sapiens]	0.17
		Homo sapiens cDNA FLJ12813 fis, clone NT2RP2002503, weakly	
4344	AK022875	similar to ZINC FINGER PROTEIN 45	1.7
4245	D10010	Arabidopsis thaliana Atpk7 gene for serine/threonine protein kinase,	
4345	D10910	complete cds	0.61

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
4346	X53096	S.aureus genes encoding Sau96I DNA methyltransferase and Sau96I restriction endonuclease	0.2
4347	M57855	Human protein S-beta pseudogene, exons 5 and 6	0.2
4348	Z73523	S.cerevisiae chromosome XVI reading frame ORF YPL167c	0.065
4349	U67562	Methanococcus jannaschii section 104 of 150 of the complete genome	0.54
4350	Y09047	M musculus pxF gene, strain 129VJ	0.002
4351	XM_027332	Homo sapiens ribosomal protein L36 (RPL36), mRNA	0.0001
4352	AF106029	Thomomys mazama clone 12-8 microsatellite sequence	0.25
4353	AL137428	Homo sapiens mRNA; cDNA DKFZp761N1323 (from clone DKFZp761N1323)	0.028
4354	L81915	Homo sapiens (subclone 1_g8 from BAC H76) DNA sequence, complete sequence	0,63
4355	J01291	soybean 7s seed storage protein alpha subunit mrna (b), from gmc- alpha'-alpha 236	3,7
4356	NM_006649	Homo sapiens serologically defined colon cancer antigen 16 (SDCCAG16), mRNA	0.69
4357	AL137635	Homo sapiens mRNA; cDNA DKFZp434E175 (from clone DKFZp434E175)	6.1
4358	Y14422	M.musculus DNA for retinal protein	0.23
4359	D32056	Human gene for 2-oxoglutarate deliydrogenase, exon 1 sequence	6.4
4360	D14337	Yeast pck1+ gene for protein kinase, complete cds	0.085
4361	J02756	Mouse band 3 anion exchange protein gene, complete cds	1.9
4362	XM_048747	Homo sapiens KIAA1223 protein (KIAA1223), mRNA	0.69
		Saccharomyces douglasii mitochondrial cytochrome c oxidase subunit	
4363	M97514	I (COXI) gene, complete cds	0.19
4364	D10910	Arabidopsis thaliana Atpk7 gene for serine/threonine protein kinase, complete cds	0.53
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
4365	AK015420	library, clone:4930448I18, full insert sequence	0.003
4366	U82487	Branchiostoma floridae engrailed protein (AmphiEn) mRNA, complete cds	0,22
4367	AB052798	Feline foamy virus env gene for envelope protein, complete cds, clone:HFFV	0.08
4368	AY027174	Arabidopsis thaliana sequence flanking 3' end of Ds-GeneTrap insertion from line GT2850	0.25
4369	AJ298684	Phylloxera sp. MBLM2 mitochondrial ATP6 gene for ATP synthase A chain subunit 6	0.67
4370	NM_013696	Mus musculus thyrotropin releasing hormone receptor (Trhr), mRNA	0.000003
4371	AF176830	Dictyostelium discoideum polyphosphate kinase (PPK) gene, complete cds	0.83
4372	AB036665	Bacteriophage WO genes, ISW1 inserted sequence, partial and complete cds	0.17
4373		Prevotella albensis dpp-IV gene for dipeptidyl peptidase IV	1.6
4374	L77040	Homo sapiens (subclone 8 c11 from P1 H22) DNA sequence	0.025

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
4400	XM_005318	Homo sapiens hypothetical protein FLJ11125 (FLJ11125), mRNA	0.44
4401	U76670	Arabidopsis thaliana recA-like protein (AtDMC1) gene, complete eds	0.051
4402	AK009448	Mus musculus adult male tongue cDNA, RIKEN full-length enriched library, clone:2310021K08, full insert sequence	1.6
4403	AY003917	Phytophthora infestans isolate 94-52 NADH dehydrogenase subunit 4	
4404		gene, partial cds; mitochondrial gene for mitochondrial product	0.16
4404	AC084160	Caenorhabditis elegans cosmid Y73B3B, complete sequence	0.21
4405	AF330120	Homo sapiens voltage-gated sodium channel type III alpha subunit (SCN3A) gene, exon 4	2.4
4406	AK022135	Homo sapiens cDNA FLJ12073 fis, clone HEMBB1002387	0.49
4407	AF332583	Homo sapiens stratum corneum chymotryptic enzyme (KLK7) gene, complete cds, alternative transcripts	1.7
4408	XM_043049	Homo sapiens hypothetical gene supported by AL080215 (LOC92138), mRNA	0.7
4409	AK017682	Mus musculus 8 days embryo cDNA, RIKEN full-length enriched library, clone:5730465109, full insert sequence	2.1
4410	AJ409162	Solanum ochranthum LINE retrotransposon within partial ure gene for urease, exons 6-8	0.025
4411	AF183584	Papio hamadryas tyrosinase gene, exon 2	0.24
4412	AK010138	Mus musculus adult male tongue cDNA, RIKEN full-length enriched library, clone:2310069P03, full insert sequence	0.56
4413	AF112141	Oryzias latipes homeodomain transcription factor (eng2) mRNA, complete cds	0,021
4414	AK022458	Homo sapiens cDNA FLJ12396 fis, clone MAMMA1002758	4.9
4415	AK024159	Homo sapiens cDNA FLJ14097 fis, clone MAMMA1000760	0.00007
		Macaca fascicularis brain cDNA clone:QtrA-10686, full insert	
4416	AB060834	sequence	2.1
4417	AF063864	Schizosaccharomyces pombe essential nuclear protein Mcm3p (mcm3+) gene, complete cds	0.21
		Drosophila melanogaster WDS (wds) and egghead (egh) genes,	0,21
4418	AF233288	complete cds	1.3
4419	AE003106	Drosophila melanogaster genomic scaffold 142000013386018, complete sequence	0.007
4421		Oryza sativa DNA for phospholipase D, complete cds	0.007
4422	AF216290	Mus musculus heterochromatin protein 1 alpha mRNA, complete cds	0.66
4423		Homo sapiens Tax interaction protein 43 mRNA, partial cds	0.18
4424	AF011446	Mus musculus granzyme K gene, complete cds	1.9
		Ceratosolen bisulcatus cytochrome oxidase subunit I (COI) gene, partial cds; tRNA-Leu gene, complete sequence; and cytochrome	
4425		oxidase subunit II (COII) gene gene, partial cds; mitochondrial genes for mitochondrial products	0.68

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1		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Zygoballus rufipes NADH dehydrogenase subunit I (ND1) gene,	1 11111111
4426	AF328031	partial cds; mitochondrial gene for mitochondrial product	1.7
		Alouatta stramineus cytochrome b gene, partial cds; mitochondrial	
4427	AF289983	gene for mitochondrial product	0.72
4428	Z73253	S.cerevisiae chromosome XII reading frame ORF YLR081w	0.19
		Homo sapiens hypothetical protein DKFZp762M115 (KIAA1630),	
4429	NM_018706	mRNA	1E-48
		Dictyostelium discoideum non-LTR retrotransposon TRE5-B,	
4430	AF298209	polyprotein (gag) and group-specific antigen (pol) genes, complete cds	0.74
4431	AJ276991	Mus musculus partial mRNA for ClpX protein	0.079
4400	4 Tho 4 ma 4 o	Human endogenous retrovirus HERV-K(II) DNA, complete sequence	
4432	AB047240	and flanking region	0.13
4433	L40389	Candida glabrata ERG11 gene, complete cds	0.027
4434	7500500	Mouse Ins DNA; cellular component that mediates integration and	
4434	K02592	excision of polyoma virus DNA	0.38
4435	D21701	Crocks (cases holy) -T-TDD C TATA 1 1: 1:	
4433	D31782	Snake (green habu) gTgTBP gene for TATA-box binding protein	1,5
4436	AK019649	Mus musculus adult male testis cDNA, RIKEN full-length enriched library, clone:4930480I16, full insert sequence	
4437	U36777	Carassius auratus aldolase C mRNA, complete cds	0.25
4438	AF036690	Caenorhabditis elegans cosmid F09C11	0.094
7730	711 030030	Cacitornabuttis cicgans cosmid 109C11	0.72
4439	NM 012750	Rattus norvegicus Tyrosine kinase receptor ligand 2 (Retl2), mRNA	0.63
4440	M74000	Brugia malayi myosin heavy chain gene, complete cds	0.073
4441	XM 042656	Homo sapiens LOC87184 (LOC87184), mRNA	0.66
		(,), and a	0.00
		Kallichroma tethys isopenicillin N synthase (pcbC) and alpha-	
4442	AF335329	aminoadipyl-cysteinyl-valine synthetase (pcbAB) genes, complete cds	6.1
		Homo sapiens region containing hypothetical protein FLJ10549;	
4443	XM_037269	hypothetical protein FLJ10517 (LOC82252), mRNA	0.26
4444	AF130079	Homo sapiens clone FLC0578 PRO2852 mRNA, complete cds	0.24
4445	AF086241	Homo sapiens full length insert cDNA clone ZD29F04	0.51
ľ		Schedorhinotermes sarawakensis 16S mitochondrial ribosomal RNA	
4446	AF262583	gene, partial sequence; mitochondrial gene for mitochondrial product	4.6
4447	AF170550	Arabidopsis thaliana germin-like protein 7 (GLP7) gene, complete cds	2
4448	AC084500	Caenorhabditis briggsae cosmid G13G15, complete sequence	0.66
4449	M23360	Trypanosoma brucei cytochrome C mRNA, partial cds	2.1
		Schizosaccharomyces pombe zinc finger protein Grt1 (grt1) gene,	
4450	AF236387	complete cds	0.076
4453	XM_010610	Homo sapiens hypothetical protein (FLJ10287), mRNA	0,027
	A3/021002	Oryza sativa microsatellite MRG4233 containing (TA)X42, closest to	
4454	AY021908	marker C563, genomic sequence	0.025
4455	AK021675	Homo sapiens cDNA FLJ11613 fis, clone HEMBA1004012	0.073

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#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE 4456 NM 030995 Rattus norvegicus Microtubule-associated protein 1a (Mapla), mRNA 1.8 4457 AF015298 Human herpesvirus 6 IE-2 protein (ie2hom) gene, complete cds 0.058 Homo sapiens hypothetical protein FLB3342 (FLB3342), mRNA 4458 XM 011297 0.0000009 4459 AE006714 Sulfolobus solfataricus section 73 of 272 of the complete genome 0.002 Plasmodium falciparum chromosome 2, section 71 of 73 of the 4460 AE001434 complete sequence 0.004 Rachiplusia ou nuclear polyhedrosis virus protein-tyrosine phosphatase (ptp), late expression factor-2 (lef-2), occlusion matrix protein (polh), capsid-associated protein, and protein kinase (pk1) 4461 AF068270 genes, complete cds; and unknown genes 0.027 Mus musculus adult male testis cDNA, RIKEN full-length enriched 4462 AK015582 library, clone:4930477O15, full insert sequence 0.08 4463 XM 039778 Homo sapiens HSPC047 protein (HSPC047), mRNA putative polymerase...capsid protein [potato aucuba mosaic potexvirus 4464 S73580 PAMV, Genomic RNA Complete, 6 genes, 7059 ntl 0.071 4465 XM 016858 Homo sapiens LOC87857 (LOC87857), mRNA 3E-13 Vibrio cholerae chromosome II, section 84 of 93 of the complete 4466 AE004427 chromosome 0.69 4467 AK025346 Homo sapiens cDNA: FLJ21693 fis. clone COL09609 4.9 4468 U89926 Drosophila melanogaster cut gene, partial sequence 4469 X57171 D.carvophyllus CARSR12 gene 2 Mus Musculus basic domain/leucine zipper transcription factor 4470 L36434 mRNA, 3' end of cds 2 Arabidopsis thaliana cultivar Landsberg erecta Arac2 (Arac2) gene, 4471 AF115469 complete cds 1.9 Petunia x hybrida taz1 gene for tapetum-specific zinc finger protein 1, 4472 AB063169 promoter region and partial cds 0.64 Bos taurus (clones L6, C15, C12, C8, C9, C17, C19, C12, C18, C5, 4473 L41691 C6, C3, C13, C10, C47) mRNA, 3' end of cds 0.47 4474 U61463 Human myosin VIIa (MYO7A) gene, 5' exon 36 0.0007 Mus musculus ATP-binding cassette, sub-family D (ALD), member 4 4475 NM 008992 (Abcd4), mRNA 1.9 4476 XM 050942 Homo sapiens ribosomal protein L6 (RPL6), mRNA 0.00009 Mus musculus adult male cerebellum cDNA, RIKEN full-length 4477 AK005146 enriched library, clone:1500004F14, full insert sequence 0.71 Hepatitis C virus isolate SCpreSC2c14 nonfunctional polyprotein 4478 AF345089 gene, partial sequence 0.71 Mus musculus adult male stomach cDNA, RIKEN full-length 4479 AK008812 enriched library, clone:2210403E17, full insert sequence 6.4 4480 XM 006983 Homo sapiens A kinase (PRKA) anchor protein 3 (AKAP3), mRNA 1.6 4481 X06438 Soybean leghaemoglobin gene lba promoter region 0.19 4482 AF020051 Blackcurrant reversion virus RNA2 polyprotein mRNA, complete cds 0.71

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	-
SEQ ID		Treatest Progness (Diastry Vs. Gendank)	
NO	ACCESSN	DESCRIP	P VALUE
4483	XM_051560		0.6
4484	AB023481	Cyprinus carpio mRNA for mitogen activated protein (MAP) kinase p38, complete cds	6.3
4485	AJ250933	Homo sapiens partial SIP1 gene for survival interacting protein 1, exons 3-4	2.2
4486	XM_039989	Homo sapiens microfibrillar-associated protein 3 (MFAP3), mRNA	0.69
4487	XM_047554	Homo sapiens similar to KIAA1473 protein (H. sapiens) (LOC92830), mRNA	6.7
4488	AB042411	Homo sapiens strg gene for striatum-specific G potein-coupled receptor, complete cds	2
4489	XM_035221	Homo sapiens ancient ubiquitous protein 1 (AUP1), mRNA	0.16
4490	D14061	Schizosaccharomyces pombe mRNA for ORF	0.015
4491	AE001426	Plasmodium falciparum chromosome 2, section 63 of 73 of the complete sequence	0,074
4492	M18349	Rat leukocyte common antigen (L-CA) gene, exons 1 through 5	0.16
4493	AF020726	Tetrahymena thermophila phosphoglucomutase (PGMI) gene, complete cds	0.058
4494	M80241	Caenorhabditis elegans unc-6 gene, complete cds	2.1
4495	U67234	Human clone HS5.29 Alu-Ya5 sequence	0.083
4496	AK000864	Homo sapiens cDNA FLJ10002 fis, clone HEMBA1000046	0.00001
4498	AK024393	Homo sapiens cDNA FLJ14331 fis, clone PLACE4000320	0.064
4499	AF071754	Mus musculus DNA cytosine methyltransferase mRNA	0.004
4500	XM_047325	Homo sapiens hypothetical protein LOC57187 (LOC57187); mRNA	0
4501	XM_043434	Homo sapiens villin-like (VILL), mRNA	1.8
4502	NM_032675	Homo sapiens hypothetical protein MGC10954 (MGC10954), mRNA	0.0008
4503	AL589203	Human DNA sequence from clone RP11-391F23 on chromosome 6, complete sequence [Homo sapiens]	0.0003
4504	M86544	Cow prostaglandin F synthetase II (PGFSII) mRNA, complete cds	0.66
4505	AF339782	Homo sapiens clone IMAGE:1871856, mRNA sequence	0
4506	Z74196	S.cerevisiae chromosome IV reading frame ORF YDL148c	6.2
4507	AF012751	Danio rerio olfactory receptor protein 2.4 mRNA, complete cds	6.8
4508	AF184796	Arabidopsis thaliana inhibitor tagged site ITS29 genomic sequence	1
4509	AE006187	Pastcurella multocida PM70 section 154 of 204 of the complete genome	1.0
4510	AE006187 AJ272171		1.8
4511	AK024656	Podospora anscrina degenerate gypsy-like retrotransposon Yeti Homo sapiens cDNA: FLJ21003 fis, clone CAE03685	0.074
4512		Lithobius forficatus mitochondrion, complete genome	1E-90 0.066
7512	110_002029	M.fascicularis mRNA for metalloprotease-like, disintegrin-like	0.066
4513	X87205	protein, IVa /	4.9

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		Andre 312 Predicts Preignoof (Blastin Vs. Genoank)	T
NO	ACCESSN	DESCRIP	P VALUE
4514	XM_011595		5E-47
		Homo sapiens cDNA FLJ11475 fis, clone HEMBA1001734,	
4515	AK021537	moderately similar to CADHERIN-11 PRECURSOR	0.068
4516	AB044877	Equus caballus DNA, microsatellite TKY377	0.076
4517 4518	AC079391	Homo sapiens clone CTD-2243M11, complete sequence	0.00001
4518	AK000932	Homo sapiens cDNA FLJ10070 fis, clone HEMBA1001581	0
4519	AE001376	Plasmodium falciparum chromosome 2, section 13 of 73 of the	
4519	XM 040095	complete sequence	0.24
4521	AF224317	Homo sapiens hypothetical protein (DKFZP434G0310), mRNA	1.8
4522	AE001155	Gallus gallus Frizzled-7 (cFz-7) mRNA, complete cds	1.5
4523	AJ313130	Borrelia burgdorferi (section 41 of 70) of the complete genome	2
4323	AJ313130	Medicago truncatula transposon (CACTA type), clone 65K06	2
4524	AF020187	Amblyomma americanum ecdsyteroid receptor (AamEcRA1) mRNA, complete cds	6,3
		Homo sapiens genomic DNA, chromosome 21q22.1, D21S226-AML	0.3
4525	AP000249	region, clone:B762O15, complete sequence	2E-19
4526	AF218380	Mus musculus CBLN3 (Cbln3) gene, complete cds	0.69
		gent, complete our	0.07
4527	AP000601	Arabidopsis thaliana genomic DNA, chromosome 3, P1 clone: MJL14	0.029
4528	AE001140	Borrelia burgdorferi (section 26 of 70) of the complete genome	0.64
		Human DNA sequence from clone RP1-317N9 on chromosome 6,	
4529	AL358852	complete sequence [Homo sapiens]	2.2
4530	NM_024596	Homo sapiens hypothetical protein FLJ12847 (FLJ12847), mRNA	0.82
4531	X12918	Human Na+,K+ ATPase gene part of intron XVI (alpha III isoform)	0.0003
4532	AF298180	Caenorhabditis elegans tropomyosin isoform IV mRNA, complete cds	0.57
4533	M21590	M.mycoides capri Gly-tRNA gene	0.48
4534	AF250227	Homo sapiens adenylyl cyclase type VI gene, partial cds	0.19
4535	Z36061	S.cerevisiae chromosome II reading frame ORF YBR192w	0.071
ľ		Endogone pisiformis 18S ribosomal RNA gene, partial sequence,	
- 1		internal transcribed spacer 1, 5.8S ribosomal RNA gene and internal	
4536	AF006510	transcribed spacer 2, complete sequence, and 26S ribosomal RNA gene, partial sequence	
4330	AF000310	gene, partial sequence	0.08
4537	D88262	Pisum sativum PsCHS5 gene for chalcone synthase, complete cds	0.075
4538	T 21504	Neurospora crassa cytoplasmic dynein heavy chain (ro-1) gene,	
4338	L31504	complete cds	2.3
4539	S81605	S100 beta =neural calcium binding protein beta subunit {5' region}	
4539		[rats, Genomic, 2279 nt]	0.66
7540	A332/3	P.falciparum complete gene map of plastid-like DNA (IR-A)	0.23
4541	U18868	Squalus acanthias multifunctional protein CAD mRNA, complete cds	0.024

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#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE 4542 AF184835 Arabidopsis thaliana inhibitor tagged site ITS68 genomic sequence 0.71 Human DNA sequence from clone RP11-509D8 on chromosome 9, 4543 AL513350 complete sequence [Homo sapiens] 0.64 Caenorhabditis elegans cosmid F48C5, complete sequence 4544 Z68107 4545 AK002184 Homo sapiens cDNA FLJ11322 fis, clone PLACE1010329 U00025 Caenorhabditis elegans cosmid PAR2, complete sequence 4546 0.46 4547 U39548 Meloidogyne artiellia mtcuticline-1 gene, 5' upstream region 0.64 4548 AK024271 Homo sapiens cDNA FLJ14209 fis, clone NT2RP3003346 0.0003 Arabidopsis griffithiana chalcone synthase gene, promoter region and 4549 AF248989 0.3 Monoclea gottschei subsp. gottschei tRNA-Thr/tRNA-Leu intergenic 4550 AF081208 spacer, partial chloroplast semence 0.009 4551 XM 002770 Homo sapiens hypothetical protein (FLJ10996), mRNA 5.8 Mus musculus MHC class I related protein 1 (MR1) gene, complete 4552 AF035672 1.1 Macrobrachium intermedium 16S ribosomal RNA gene, partial 4553 AF374466 sequence; mitochondrial gene for mitochondrial product 5.9 pdm-2=POU domain gene [Drosophila melanogaster, mRNA, 2171 4554 S80559 0.009 Homo sapiens aminophospholipid-transporting ATPase (ATP10C) 4555 AY029489 gene, exon 3 1E-89 Mus musculus RIKEN cDNA 4930433D19 gene (4930433D19Rik). 4556 NM 026255 2.2 Homo sapiens partial GDH gene for uridine diphospho-glucose 4557 AI276163 dehydrogenase, exons 8-11 0.0002 4558 NC 001566 Apis mellifera ligustica mitochondrion, complete genome 0.19 Mus musculus, calreticulin, clone MGC:6209 IMAGE:2655918. 4559 BC003453 mRNA, complete cds 2 XM_048784 Homo sapiens mature T-cell proliferation 1 (MTCP1), mRNA 4560 0.62 4561 AF119845 Homo sapiens PRO1304 mRNA, complete cds 0.01 4562 AF251477 Plasmodium berghei glutathione reductase gene, partial crls Homo sapiens ATPase, Na+/K+ transporting, beta 3 polypeptide 4563 XM 018443 (ATP1B3), mRNA 2 Trichomonas vaginalis calcium motive P-type ATPase (CA-2) gene. 4564 AF145282 nartial cds 0.11 4565 AF280548 Homo sapiens neuropilin-1 (NRP1) gene, exons 1, 2, and partial cds 2.1 4566 NC 002253 Buchnera sp. APS plasmid pLeu, complete sequence 0.24 4567 AJ238613 Gallus gallus mRNA for teneurin-1 0.61 4568 Y08926 P.falciparum mRNA for AARP1 protein, partial 0.21 4569 Z86105 A.thermophilum celA gene and manA pseudogene 0.69 S.pombe chromosome I p1 p32A8 4570 AL583902 1.1 Human DNA sequence from clone RP3-456L16 on chromosome 6. 4571 complete sequence [Homo sapiens] AL358951 0.072 4572 AJ001164 Drosophila melanogaster additional sex combs cDNA sequence 0.1

SEQ ID NO   ACCESSN   Schistosoma mansoni thioredoxin peroxidase 3 (TPx3) gene, complete des   0.19				
ACCESSN			Table 3A Nearest Neighbor (BlastN vs. Genbank)	
AF301004   Schistosoma mansoni thioredoxin peroxidase 3 (TPx3) gene, complete clds   0.19				T
AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004   AF301004	NO	ACCESSN		P VALUE
4573			Schistosoma mansoni thioredoxin peroxidase 3 (TPx3) gene, complete	· ·······
AF276995   Schopus laevis homeobox protein Six4.2 (Six4.2) gene, partial cds   2	4573	AF301004		0.19
4576				V.12
4576	4574	AF276995	Xenopus laevis homeobox protein Six4.2 (Six4.2) gene, partial cds	,
Human DNA sequence from clone RP11-76G11 on chromosome X, complete sequence [Homo sapiens]	4575	BC007465	Homo sapiens, clone IMAGE:3828547, mRNA	
4576   AL589824   complete sequence [Homo sapiens]   0.17     4577				0.00002
AF151171   Gritthorlhynchus anatimus mannose 6-phosphate/insulin-like growth factor 2 receptor (M6p/lg/2r) gene, exons 2 and 3 and partial eds   0.67	4576	AL589824		0.17
4577				0.17
4577	1 1		Ornithorhynchus anatimus mannose 6-phosphate/insulin-like growth	
4578	4577	AF151171	factor 2 receptor (M6p/Igf2r) gene, exons 2 and 3 and partial cds	0.67
4590   MM_016609   Homo sapiens KIAA1046 protein (KIAA1046), mRNA   1.1	4578	AK000883	Homo sapiens cDNA FLJ10021 fis clone HEMBA 1000568	
Home sapiens transcriptional intermediary factor 1 gamma   2	4579	XM 016630	Homo sapiens KIA A 1046 protein (KIA A 1046) mPNA	
4580   NM_015906   CIFFIGAMMA), transcript variant alpha, mRNA   2			Homo saniens transcriptional intermediary factor 1 gamma	1.1
AF274053   Xenopus laevis maxi-K potassium channel alpha subunit Slo mRNA, complete cds   Pseudoregma pendleburyi mitochondrial genes for small subunit fNNA, (RNA-Val, large subunit rRNA, partial and complete   Sequences   0.003   Af2003874   Sequences   0.003   Af2003874   Sequences   0.003   Af2003874   Sequences   0.003   Af2003874   Sequences   0.004   Af203874   AF277081   protein 1 (bys1) gene, complete cds   0.26   Homo sapiens chromosome V landmark: proximal external boundary   0.69   AF34545   AF277081   Pyodit yiPL6 gene for ATPase   0.23   AF27001   Ebola virus subtype Zaite strain Mayinga complete genome   1.5   AF272001   Ebola virus subtype Zaite strain Mayinga complete genome   1.5   AF27201   Ebola virus subtype Zaite strain Mayinga complete genome   1.5   AF27201   Ebola virus subtype Zaite strain Mayinga complete genome   1.5   AF27201   Ebola virus subtype Zaite strain Mayinga complete genome   1.5   AF27201   Ebola virus subtype Zaite strain Mayinga complete genome   1.9   AF27201   Ebola virus subtype Zaite strain Mayinga complete genome   1.9   AF27201   Ebola virus subtype Zaite strain Mayinga complete genome   1.9   AF27201   Ebola virus subtype Zaite strain Mayinga complete genome   1.9   AF27201   A	4580	NM 015906	(TIF1GAMMA) transcript variant alpho mRNA	2
ASS				
Pseudoregma pendlebuyi mitochondrial genes for small subunit rRNA, IRNA-Val, large subunit rRNA, partial and complete sequences   0.003	4581	AF274053		0.10
AB035874   Sequences   AB035874   Sequences   AB035874   Sequences   AB035874   Sequences   AB035874   Sequences   AJellomyces dermatitidis strain 104 blastomyces yeast phase-specific protein 1 (bys1) gene, complete cds   O.26   Homo sapiens chromosome V landmark: proximal external boundary of P7 VCV inverted repeat   O.59   AF585   AF343561   AF343561   AF343561   AF34561   AF34565   AF34561   AF3456	1001	111271000		0.19
4583			rPNA tPNA-Val large cultural rPNA montal and assument	
Ajellomyces dermatitidis strain 104 blastomyces yeast phase-specific protein 1 (bys1) gene, complete ods   0.26	4582	A B025974		
4584   AF37081   Introduction   Internation   Introduction   Introduction   Introduction   Introduction   Introduction   Introduction   Introduction   Internation   Inter	4302	AD033674		0.003
Homo sapiens chromosome Y landmark: proximal external boundary   0.69	4583	A E277081		
4584   AF334561   Of P7 VCY inverted repeat   0.69	4303	AL277001		0,26
4585   X55197   P.yoelii YEL6 gene for ATPase   0.03	4584	A 1722/1561		0.00
4586   L32141   Mitochondrion Lymantria dispar ribosomal RNA large sabunit   0.008   4587   AF272001   Ebola virus subtype Zaire strain Mayinga complete genome   1.5   Vibrio cholerae chromosome II, section 62 of 93 of the complete   4588   AE004405   chromosome   Home sapiens debranching enzyme (S. Cerevisiae) homolog 1   4589   XM_010966   (DRB1), mRNA   2   4591   U68246   Dictyostelium discoideum WacA (wacA) gene, complete cds   0.002   4592   XM_047617   Home sapiens KIAA1349 protein (KIAA1349), mRNA   0.027   4593   AK003428   dibrary, clone:111000407, full insert sequence   2.2   4594   XM_029072   Home sapiens LOS89256 (LOC89256), mRNA   0.009    Buchnera aphidicola aspartyl-RNA synthesae (aspS), thioredoxin reductase (trxB), seryl-tRNA synthesae (serS), phosphoserine aminotransferase (serC), 3-enolpyruvylshikimate-5-phosphate synthesae (asroA), ribosomal protein S1 (rpsA), integration host fa- 0.21   4596   AK023907   Home sapiens GNNA FILJ3484 fis, chom THYRO1000815   0   4598   AB042649   Maternation and the strain of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constraints of the same constr				
4587   AF272001   Ebola virus subtype Zaire strain Mayinga complete genome   1.5				
AB004405   Chromosome   1.9			Ehola vivus cultura Zaira strain Maria sa associat RNA large subunit	
4588	4367	At 2/2001	Wibrio chalenge characters at a series complete genome	1.5
Homo sapiens debranching enzyme (S. Cerevisiae) homolog 1   2	4500	AE004405		
ASS   XM_019966   (DBR1), mRNA   2	4300	, AE004403		1.9
4591   U68246   Dictyostellum discoideum WacA (wacA) gene, complete eds   0.002	4500	3/3 / 0100//		
4592 XM_047617   Homo sapiens KIAA1349 protein (KIAA1349), mRNNA   0.027				
Mus musculus 18 days embryo cDNA, RIKEN full-length enriched   2.2			Dictyostellum discordeum wacA (wacA) gene, complete cds	
AB003428   Library, clone:111000407, full insert sequence   2.2	4392	XM_04/61/		0.027
ASM   029072   Homo sapiens LOC89256 (LOC89256), mRNA   0.009	4502	177000100	Mus musculus 18 days embryo cDNA, RIKEN full-length enriched	
Buchnera aphidicola aspartyl-fRNA synthease (aspS), thioredoxin reductase (trxB), seryl-tRNA synthase (serS), phosphoserine aminotransferase (serC), 3-enolpyruvylshikimate-5-phosphate synthetase (asrA), ribosomal protein S1 (rpsA), integration host fa- 4596 AK023907 Homo saptions GDNA FL13484 fis, chone THYRO1008015 0 Watermelon spotted wilt virus nss. np genes for non-structural, mucleocapsid protein, complete cds, isolate:WS-O 0.024 Bacillus sp. DNA for cytochrone c oxidase subunit I,II and III,			library, clone:1110004J07, full insert sequence	
roductase (trxB), seryl-tRNA synthase (serS), phosphoserine aminotransferase (serC), 3-enolypruvylshikimates-phosphate synthetase (aroA), ribosomal protein S1 (rpsA), integration host fa> 0.21	4394	XM_029072	Homo sapiens LOC89256 (LOC89256), mRNA	0.009
roductase (trxB), seryl-tRNA synthase (serS), phosphoserine aminotransferase (serC), 3-enolypruvylshikimates-phosphate synthetase (aroA), ribosomal protein S1 (rpsA), integration host fa> 0.21				1
AB042649	1		Buchnera aphidicola aspartyl-tRNA synthetase (aspS), thioredoxin	- 1
4595         L43549         synthetase (aroA), ribosomal protein S1 (rpsA), integration host fa>         0.21           4596         AK023907         Homo sapiens cDNA FLJ13845 fis, clone THYRO1000815         0           4598         AB042649         Watermelon spotted wilt virus nss, np genes for non-structural, uncleocapsid protein, complete cds, isolate:WS-O         0.024           Bacillus sp. DNA for cytochrome c oxidase subunit I,II and III,         0.024			reductase (trxB), seryl-tRNA synthase (serS), phosphoserine	
4596         AK023907         Homo sapions cDNA FLJ13845 fis, clone THYRO1000815         0           Watermelon spotted will virus nss, np genes for non-structural, and clocoapsid protein, complete cds, isolate:WS-O         0,024           Bacillus sp. DNA for cytochrome c oxidase subunit I,II and III,         and III,			aminotransferase (serC), 3-enolpyruvylshikimate-5-phosphate	
Watermelon spotted wilt virus nss, np genes for non-structural, mucleocapsid protein, complete cds, isolate:WS-O 0.024 Bacillus sp. DNA for cytochrome c oxidase subunit I,II and III,			synthetase (aroA), ribosomal protein S1 (rpsA), integration host fa>	0.21
4598 AB042649 nucleocapsid protein, complete cds, isolate:WS-O 0.024  Bacillus sp. DNA for cytochrome c oxidase subunit I,II and III,	4596	AK023907	Homo sapiens cDNA FLJ13845 fis, clone THYRO1000815	0
Bacillus sp. DNA for cytochrome c oxidase subunit I,II and III,			Watermelon spotted wilt virus nss, np genes for non-structural,	
	4598	AB042649	nucleocapsid protein, complete cds, isolate:WS-O	0.024
4599 D85547 complete and partial cds 0.23				
	4599	D85547	complete and partial cds	0.23

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#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEO ID NO ACCESSN DESCRIP P VALUE Human (clone D13-2) L-iditol-2 dehydrogenase gene, exon 4, exon 5, 4600 L29252 exon 6 and exon 7 2E-08 Xylella fastidiosa 9a5c, section 18 of 229 of the complete genome 4601 AE003872 1.5 4602 U29924 Human AMP deaminase (AMPD3) gene, exon 12 0.0003 4603 AE000653 Helicobacter pylori 26695 section 131 of 134 of the complete genome 0.75 4605 AP001235 Homo sapiens genomic DNA, chromosome 2p11.2, clone:cos607/7B8 0.024 4606 AF207725 Mustela lutreola isolate D23 mitochondrial D-loop, partial sequence 0.028 4607 AE007377 Streptococcus pneumoniae section 60 of 194 of the complete genome 0.083 4608 XM 018430 Homo sapiens LOC88330 (LOC88330), mRNA 4E-27 Plasmodium falciparum DNA-dependent RNA polymerase (TRNAP) 4609 AF273674 gene, complete cds; nuclear gene for probable mitochondrial product 0.002 Oryza sativa microsatellite MRG1244 containing (AT)X23, genomic AY018919 4610 sequence 6.7 Arabidopsis thaliana genomic DNA, chromosome 5, BAC AB025642 4611 clone:T32G24 2.1 4612 X59275 D.melanogaster mRNA for posterior sex combs 0.68 4613 AF052959 Homo sapiens type XV collagen (COL15A1) gene, exon 6 0.71 4614 U32730 Haemophilus influenzae Rd section 45 of 163 of the complete genome 0.025 Bolitoglossa mexicana isolate MEX_6C cytochrome b gene, partial 4615 AF212976 cds; mitochondrial gene for mitochondrial product 0.076 Cancer magister molt-inhibiting hormone precursor mRNA, complete 4616 AF031493 0.22 4617 AJ245965 Danio rerio mRNA for pbxy homeodomain protein (pbxy gene) 0.068 4618 XM 031524 Homo sapiens AF15q14 protein (AF15Q14), mRNA 0 4619

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SEO TO		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUI
4628	X55978	T. brucei ESAG 8 gene for a leucine-rich repeat family protein	0.24
4629	U41529	Caenorhabditis elegans cosmid EGAP3	0.59
		Homo sapiens associated molecule with the SH3 domain of STAM	0.39
4630	XM_039502	(AMSH), mRNA	5.3
4631	AF141650	Nyctalus noctula microsatellite P223 sequence	1.5
		Human DNA sequence from clone RP6-152C18 on chromosome	
4633	AL096887	Xq26.1-27.3, complete sequence [Homo sapiens]	0.02
		Homo sapiens mRNA; cDNA DKFZp434E2221 (from clone	
4634	AL137458	DKFZp434E2221)	0.074
4635	AE002292	Chlamydia muridarum, section 24 of 85 of the complete genome	5.1
4636	XM_048200		5.4
		Dictyostelium discoideum LTR-retrotransposon Skipper, partial	
4637	AF017041	genomic sequence, 5' end	0.16
4638	Z98853	Caenorhabditis elegans cosmid R08A2, complete sequence	5.7
4639	XM_017899		0
- 1		Mus musculus, Similar to RIKEN cDNA 3230401N03 gene, clone	
4640	BC006943	MGC:6906 IMAGE:2655807, mRNA, complete cds	2
J		Lycopersicon esculentum MADS-box transcription factor jointless	
4641	AF275345	gene, complete cds	0.62
- 7		Mus musculus adult male cecum cDNA, RIKEN full-length enriched	
4642	AK018612	library, clone:9130011L11, full insert sequence	1.9
- 1		Streptococcus pyogenes M1 GAS strain SF370, section 97 of 167 of	
4643	AE006568	the complete genome	6.6
4644	AF057019	Dictyostelium discoideum interaptin (abpD) gene, complete cds	0.67
		Homo sapiens immune dysregulation, polyendocrinopathy.	
4645	XM_010198		5E-29
4646	NM_022599	Rattus norvegicus outer membrane protein (Omp25), mRNA	1E-14
- 1		HIV-1 isolate 965 clone 26 from Malawi, envelope glycoprotein (env)	
4647	U08455	gene, partial cds	0.72
4648	AK023304	Homo sapiens cDNA FLJ13242 fis, clone OVARC1000578	6
T		Gallus gallus fibrinogen alpha and alpha-E subunit gene, exons 5 and	
4649	U20803	6, and complete cds for alpha and alpha-E subunits	2
		Homo sapiens leukocyte immunoglobulin-like receptor 5 (LIR5) gene,	
4650	AF189768	complete cds	0.025
T		Mus musculus adult male kidney cDNA, RIKEN full-length enriched	
4651	AK002614	library, clone:0610012K15, full insert sequence	2.1
4652	AF010568	Plasmodium falciparum microsatellite TA80 sequence	0.05
T		Schizosaccharomyces pombe sulfide dehydrogenase (hmt2) gene,	
4653	AF042283	complete cds	0.63
4654		Daucus carota mRNA for neutral invertase	0.22
		Homo sapiens hypothetical gene supported by AF001893; AH005586	
	XM_035796	(LOC91063), mRNA	1.5
4656	AB008565	Homo sapiens gene for deoxylribonuclease II, 5'-upstream region	0.73
4657	Y17275	Lycopersicon esculentum p69a gene, complete CDS	0.51
4658	X03100	Human HLA-SB(DP) alpha gene	0.23

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	-	Table 3A. Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			Τ
NO	ACCESSN	DESCRIP	P VALUE
		Gallus gallus DNA polymerase gamma mRNA, nuclear gene	1 VILLOD
4685	U60297	encoding mitochondrial protein, partial cds	0.65
		D.melanogaster (strain Ja-F) alcohol dehydrogenase gene (allele Adh-	1 0.05
4686	M17837	F), complete cds	0.08
		Human germline kappa-immunoglobulin variable region (V-kappa-II)	
4687	X17265	gene Vcc	0.25
		Homo sapiens similar to hypothetical protein MGC5384 (H. sapiens)	
4688	XM_030896		7E-32
4689	AF083064	Mus musculus paxillin-like protein (Hic5) gene, complete cds	e-103
		L.helveticus galK, galT and galM genes for galactokinase, galactose-1	
4690	X57248	P uridyl transferase and mutarotase, partial cds	0.003
		Mus musculus integral membrane protein 2A (Itm2a) gene, complete	
4692	AF074020	cds	0.059
4693	D17668	Clostridium septicum gene for alpha-toxin, complete cds	0.08
		Molothrus badius cytochrome b (cytb) gene, mitochondrial gene	
4694	AF089042	encoding mitochondrial protein, partial cds	0.076
		Mus musculus adult male lung cDNA, RIKEN full-length enriched	
4695	AK004553	library, clone:1200002O22, full insert sequence	0.63
4505		Bacillus thuringiensis insecticidal crystal protein (NRcryV) gene,	
4696 4697	AF047579	complete cds	0.69
4097	AF049489	Canis familiaris factor VIII mRNA, complete cds	6.3
4698	3257500	S.xylosus xylR, xylA, xylB genes for Xyl repressor, xylose isomerase,	
4098	X57599	xylulokinase	0.024
4699	L04873	Mus musculus cystic fibrosis transmembrane regulator (CFTR) gene,	
4033	1040/3	exon 1, and upstream region	0.25
4700	XM 041483	Homo sapiens hypothetical protein MGC15827 (MGC15827), mRNA	
4700	201_041483	Plocamium cartilagineum complete internal transcribed spacer region	2.2
4701	U30355	including ITS 1, 5.8s rDNA and ITS 2	
-1701	030333	Gallus gallus myostatin (MSTN) gene, exons 1,2 and 3 and complete	0.082
4702	AF346599	cds	1.0
	122,000	COLD	1.8
4703	XM 007118	Homo sapiens A kinase (PRKA) anchor protein 11 (AKAP11), mRNA	6.2
-		Homo sapiens superkiller viralicidic activity 2 (S. cerevisiae homolog)	0.2
4704	XM 042395	like (SKIV2L), mRNA	3.5
		Into (044) 722); miletin	3.3
4705	Z78791	H.sapiens flow-sorted chromosome 6 HindIII fragment, SC6pA16H6	2E-08
		Lactococcus lactis subsp. lactis IL1403 section 164 of 218 of the	ZE-08
4706	AE006402	complete genome	0.7
4707	X97876	H.sapiens EP4 prostaglandin receptor pseudogene C17	6.6
$\neg$		Homo sapiens genomic DNA, chromosome 21q22.2, clone:T695,	
4708	AP001438	LB7T-ERG region, complete sequence	0.0001
		Homo sapiens WW domain binding protein 4 (formin binding protein	3.0001
4709	XM_049375	21) (WBP4), mRNA	3E-16
		Luehdorfia longicaudata mitochondrial ND5 gene for NADH	-22.0
4710	AB016828	dehydrogenase subunit 5, partial cds	0.45

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			Τ
NO	ACCESSN		P VALU
4711	U23439	Bovine herpesvirus 4 1.1-kb mRNA	0.64
4510		Homo sapiens genomic DNA, chromosome 21q22.2, clone:T695,	—— <u> </u>
4712	AP001438	LB7T-ERG region, complete sequence	0.00003
4713	170000000	Arabidopsis thaliana genomic DNA, chromosome 5, BAC	
4714	AP000378 AC064850	clone:T4M5	0.24
4715	Z93943	Homo sapiens clone RP11-117H9, complete sequence	0.009
4716	AJ132901	Human DNA sequence from cosmid U235H3 on chromosome X	0.006
4/10	AJ132901	Drosophila guanche mitochondrial A+T-rich region	0.052
4717	AE006291	Lactococcus lactis subsp. lactis IL1403 section 53 of 218 of the	
4/1/	AE000291	complete genome	0.24
4718	AF227741	Rattus norvegicus protein kinase WNK1 (WNK1) mRNA, complete eds	
4720	Z73524		6.1
4721	AF142484	S.cerevisiae chromosome XVI reading frame ORF YPL168w	0.072
4/21	AF142484	Sebastes melanops microsatellite Smc2 sequence	0.19
4722	X51938	D. melanogaster DNA for a region 3' to the white gene, corresponding	
4723	Z80361	to the FB-NOF insertion site in TE146	2
4723	2,80361	H. sapiens HLA-DRB pseudogene, repeat region;	0.55
4724	AF312620	Plasmodium vivax clone C19b thrombospondin-related adhesive protein gene, partial cds	
4724	AI-312020	protein gene, partial cus	2.1
4725	XM_017198	Homo sapiens hypothetical protein FLJ12085 (FLJ12085), mRNA	1E-35
		Homo sapiens chromosomal breakpoint fragment derivative 11,	1L-33
4726	AJ408943	MLL/AF4 fusion, patient UPN051	0.001
4727	XM_050754	Homo sapiens KIAA1594 protein (KIAA1594), mRNA	0.5
		Drosophila melanogaster genomic scaffold 142000013385520,	0.5
4728	AE002938	complete sequence	0.008
ĺ		Plasmodium falciparum chromosome 2, section 27 of 73 of the	
4729	AE001390	complete sequence	0.21
4730	AK000109	Homo sapiens cDNA FLJ20102 fis, clone COL04783	2.1
		Trichoderma harzianum mitochondrial plasmid pThr1, complete	
4731	AF163325	plasmid sequence	0.028
		Homo sapiens mRNA; cDNA DKFZp434M1114 (from clone	
4732	AL137277	DKFZp434M1114)	1.9
4733	T742145	· · · · · · · · · · · · · · · · · · ·	
4/33	U43145	Plasmodium chabaudi repeat organellar protein gene, complete cds	0.64
472.4	YELOES.		
1734	X71875	H. sapiens gene for protein serine kinase PSK-H1, first coding exon	1.7
1735	4 E001 450	Helicobacter pylori, strain J99 section 40 of 132 of the complete	
+/35	AE001479	genome	0.74
736	M29586	Elops saurus Ig germline H-chain pseudogene V-region, partial cds	
		Mus musculus adult retina cDNA, RIKEN full-length enriched	0,66
1737	AK020838	library, clone: A930009F24, full insert sequence	
		Melon yellow spot virus NSs and N genes for nonstructural protein	0.025
738	AB038343	and nucleoprotein, complete cds, isolate: Tospo-melo	0.00
739		Candida maltosa gene for EPD1, complete cds	0.68
		Canada minicoa Bone for Li D1, confipiete cus	0.25

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ II		Teat of Troughton (Diaset 78. Genoalik)	
NO	ACCESSN	DESCRIP	P VALUE
4740	AF361581	Arabidopsis thaliana AT4g35320/F23E12_120 gene, complete cds	1.9
4741	Z75032	S.cerevisiae chromosome XV reading frame ORF YOR124c	0,025
4742	U17681	Human neurofibromatosis 1 (NF1) gene, exons 16, 17, 18 and 19a	0.45
4743	AK025048	Homo sapiens cDNA: FLJ21395 fis, clone COL03557	0.43
4744	AF198444	Homo sapiens 10q21 mRNA sequence	0.008
		Escherichia coli O157:H7 EDL933 genome, contig 3 of 3, section 175	0.008
4745	AE005556	of 290	0.71
4746	X74784	M.musculus mk2e mRNA	0.71
4747	AB029480	Oryzias latipes gene for Y3 olfactory receptor, complete cds	0.17
	713025400	Median on policy and the Line and the Complete cuts	0.077
4749	AF072932	Medicago sativa cold acclimation responsive protein CAR1 (CAR1)	
4/49	AF0/2932	mRNA, complete cds	0.66
		Capitonus sp. 16S ribosomal RNA gene, partial sequence;	
4750	AF176056	mitochondrial gene for mitochondrial product	0.009
4751	X03366	Bovine spleen trypsin inhibitor II (SI) gene	0.12
		Human immunodeficiency virus type 1 (UG275) proviral DNA	
4752	L22951	encoding env, tat, vpu, rev, and nef genes	0.23
		Amblyomma darwini 12S ribosomal RNA gene, mitochondrial gene	
4753	U95851	for mitochondrial RNA, partial sequence	5.4
		Plasmodium falciparum chromosome 2, section 9 of 73 of the	
4754	AE001372	complete sequence	2.2
4755	AF147813	Protrama radicis elongation factor 1 alpha gene, partial cds	0.24
		Tribolium castaneum transcription factor homolog (Tc-eve) gene,	0.24
4756	U77974	complete cds	0.003
4757	XM 045423	Homo sapiens KIAA0701 protein (KIAA0701), mRNA	6.3
4758	J04357	Red clover necrotic mosaic virus RNA-1, complete sequence	
.,,,,	304357	Home continue O links Alva Alva Complete sequence	0.96
4759	<b>TR</b> 6 0404-4	Homo sapiens O-linked N-acetylglucosamine (GlcNAc) transferase (UDP-N-acetylglucosamine:polypeptide-N-acetylglucosaminyl	
4/39	XM_010136	transferase) (OGT), mRNA	0.69
4760	XM_030855	Homo sapiens similar to KIAA1361 protein (H. sapiens) (LOC90315), mRNA	0.008
		HIV-1 isolate NC5000-1999 from USA pol polyprotein (pol) gene,	
4761	AY031575	partial cds	6.2
		Homo sapiens Homer, neuronal immediate early gene, 2 (HOMER-	
4762	XM_007592	2B), mRNA	2.2
4763	AC016707	Homo sapiens BAC clone RP11-221K4 from Y, complete sequence	0.75
4764	AF001848	Oxytricha nova phosphoglycerate kinase (PGK) gene, complete cds	5.7
T	7.500.550	Mus musculus, clone MGC:7897 IMAGE:3582679, mRNA, complete	
4765	BC006693	cds	1.9
- 1		Homo sapiens (subclone 7_g1 from P1 H11) DNA sequence, complete	
4766	L81574	sequence	0.00003
4767	U67508	Methanococcus jannaschii section 50 of 150 of the complete genome	0.23

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	T :		
		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALU
	1	Oryza sativa microsatellite MRG3819 containing (TA)X27, genomic	
4768	AY021494	sequence	0.023
	1	Borrelia burgdorferi flagellar hook protein (flgE), flbD, flagellar	
		motor apparatus (motAB), fliL, fliM, fliZ, flagellar export apparatus	ľ
4769	L75945	(fliPQR, flhB), flhF, flbE genes	0.76
4770	AL590606	S.pombe chromosome II BAC pB7E8	0.068
4771	A TO CO TO O	Conospermum mitchelii chloroplast atpB-rbcL intergenic spacer	
4//1	AF060728	region, partial sequence	0.2
4772	AB056833	Macaca fascicularis brain cDNA clone:QflA-14255, full insert	
4//2	AD030633	sequence	0.068
4773	AP001416	Homo sapiens genomic DNA, chromosome 21q22.2, clone:D34, LB7T-ERG region, complete sequence	
4773	AF001410	Libra seriore analysis for the first time to the control of the co	0.0003
4774	XM 007838	Homo sapiens nuclear factor of activated T-cells 5, tonicity-responsive (NFAT5), mRNA	
4775	AC024777	Caenorhabditis elegans cosmid Y42H9AR, complete sequence	0.59
4775	110024777	Dictyostelium discoideum beta-alanine synthase (pyd3) mRNA.	0.74
4776	AF333186	complete cds	0.055
	111 555 100	Anthocidaris crassispina mRNA for dynein beta-heavy chain,	0.057
4778	D01021	complete cds	0.60
4779	U65750	Xenopus laevis fork head related (XFD-1') gene, complete cds	0.69
4780	NM 008176	Mus musculus GRO1 oncogene (Gro1), mRNA	1.8
4781	M88115	Hylobates lar Myc gene, complete cds	0.73
		, see and say of good, complete out	0.73
4782	AP000388	Arabidopsis thaliana genomic DNA, chromosome 3, P1 clone:MRI12	0.022
		Sinorhizobium meliloti plasmid pSymA section 52 of 121 of the	0,022
4783	AE007246	complete plasmid sequence	6.7
4784	XM_043069	Homo sapiens KIAA1464 protein (KIAA1464), mRNA	0.08
		Human Down Syndrome region of chromosome 21 genomic sequence,	
4785	U41482	clone A39D1-1H8	0.001
- 1		Homo sapiens mRNA; cDNA DKFZp564H072 (from clone	
4786	AL110134	DKFZp564H072)	0.058
4787	XM_005044	Homo sapiens KIAA0628 gene product (KIAA0628), mRNA	0.002
4788	XM_052430	Homo sapiens E3 ubiquitin ligase SMURF1 (SMURF1), mRNA	2E-30
		Rattus norvegicus cadherin EGF LAG seven-pass G-type receptor 3	
4790	NM_031320	(Celsr3), mRNA	1.9
4791	AF184590	Plasmodium falciparum ADA2-like protein gene, partial cds	0.009
4792	AF061434	Vicia faba amino acid transporter a (AAPA) mRNA, partial cds	1.9
4793	AL117594	Homo sapiens mRNA; cDNA DKFZp564C1563 (from clone	
		DKFZp564C1563)	0.074
4794	AJ133025	Megaselia abdita zen gene, exons 1-2	0.02
4705	VM 022005	Transaction to all all all and all all all all all all all all all al	
4795	XM_033087	Homo sapiens hypothetical protein FLJ20359 (FLJ20359), mRNA	0.024
4796	AE001462	Helicobacter pylori, strain J99 section 23 of 132 of the complete	
4/90	AE001462	genome	1.1
4797	XM 042814	Hama series benefit to the series	
+/7/	AWI_042814	Homo sapiens hypothetical protein FLJ13397 (FLJ13397), mRNA	0.063

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		Table 24 Manual III on an an	
SEQ ID		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
NO	ACCESSN	DESCRIP	1
****	NCCLUBIT	Homo sapiens mRNA; cDNA DKFZp434M0835 (from clone	P VALU
4798	AL137325	DKFZp434M0835)	25.1
		Mus musculus Chetk-alpha gene for choline/ethanolamine kinase-	2E-14
4799	AB030620	alpha, exon 4, 5, 6, 7, 8	0.64
4800	AB037417	Owene notice over 6	
4801	AK024522	Oryza sativa gene for aspartate carbamoyl transferase, complete cds Homo sapiens cDNA: FLJ20869 fis, clone ADKA02377	0.026
		The state of the s	0.21
4802	X62463	H.sapiens 3' flanking region for estrogen receptor (placenta) gene	0.7
4803	AE001127	Borrelia burgdorferi (section 13 of 70) of the complete genome	0.7
4804	Z70751	Caenorhabditis elegans cosmid F11A3, complete sequence	0.22
4805	XM_038231	Homo sapiens KIAA0793 gene product (KIAA0793), mRNA	6E-53
4806	AF223425	Mus musculus MAB21L2 (MAb21l2) gene, complete cds	4.5
		Homo sapiens hypothetical protein MGC11303 similar to Zink	
4807	XM_039048	transporter 2 (MGC11303), mRNA	2
4808	U75604	Tilapia nilotica type I estrogen receptor mRNA, complete cds	0.057
		Homo sapiens stimulated trans-acting factor (50 kDa) (STAF50),	0.057
4809	XM_015243	mRNA	1.7
		Caenorhabditis elegans cell cycle checkpoint protein Rad9 gene,	<del></del>
4810	AF247970	complete cds	0.008
$\neg$			
4811	X12574	Parsley PcPR1-2 mRNA for pathogenesis-related protein type A	0.077
4812	AF332562	Simian virus 40 strain 777, complete genome	0.64
		Homo sapiens general transcription factor IIE, polypeptide 2 (beta	
4813	XM_005118	subunit, 34kD) (GTF2E2), mRNA	1
		Mus musculus, Similar to RIKEN cDNA 2310035M22 gene, clone	
4814	BC006700	MGC:6680 IMAGE:3501285, mRNA, complete cds	0.21
		Yaba monkey tumor virus DNA, BamH1 restriction fragment E, M	
4815	AB025319	and partial C, partial and complete cds	0.18
		Homo sapiens ATPase, H+ transporting, lysosomal (vacuolar proton	
4816	XM_048433	pump), subunit 1 (ATP6S1), mRNA	0.55
4817	Y11095	Rice stripe virus RNA 3	5.9
T I		Human DNA sequence from clone RP11-241J5 on chromosome 6,	
4818	AL589991	complete sequence [Homo sapiens]	3E-78
4819	XM_043498	Homo sapiens G protein-coupled receptor 44 (GPR44), mRNA	0,0002
4820	XM_047477	Homo sapiens KIAA1638 protein (KIAA1638), mRNA	0.077
		Streptococcus thermophilus ORF1, ORF2, ORF3, ORF4, ORF5 and	
4821	AJ242480	ORF6 DNA, strain St0	0.18
		Homo sapiens region containing hypothetical protein; mutL (E. coli)	
	XM_016395	homolog 3 (LOC82389), mRNA	6E-74
4823	XM_043492	Homo sapiens 42259 (KIAA1728), mRNA	0.000000
4824	L43603	Gallus gallus Na+/K+-ATPase alpha-1 subunit gene, promoter region	0.41
		Rattus norvegicus multidrug resistance protein 1a (Pgy1) mRNA,	
4825	AF257746	complete cds	0.5
4826	X68794	D.melanogaster "ten-a" mRNA for secreted protein	1.2

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ II		greet (Saint Vis. Genbank)	T
NO	ACCESSN		P VALUE
		Homo sapiens voltage-gated sodium channel type III alpha subunit	1 VALOL
4827	AF330118	(SCN3A) gene, exon 2	0.12
4828	XM_011230		5.8
		Homo sapiens hypothetical gene supported by AL133011	
4829	XM_030669		1.9
4830	U36927	Plasmodium yoelii rhoptry protein gene, complete cds	0.21
4831	AF201353	Columba livia glutamate receptor C gene, partial cds	5
4832	AF331435	HHIVI DOC . A W	
4832	AF331433	HIV-1 D2 from Australia envelope protein (env) gene, partial cds	0.07
4833	AF362391	Candida albicans heat shock protein Hsp104 (HSP104) gene, HSP104- b allele, complete cds	
-1055	7H 302371		1.1
4834	XM_027927	Homo sapiens hypothetical gene supported by Z97017 (LOC89989), mRNA	
4835	AE001127	Borrelia burgdorferi (section 13 of 70) of the complete genome	0.48
4836	AL390126	S. pombe chromosome III cosmid c233	0.72
4837	AF110213	Rattus norvegicus insulin receptor precursor, exon 14	0.023
		Paramecium aurelia immobilization antigen (51C) gene, exon 5 and	0.56
4838	M38534	complete cds	0.015
		Plasmodium falciparum chromosome 2, section 67 of 73 of the	0.013
4839	AE001430	complete sequence	0.052
			0.032
4840	AF181475	Mus musculus synaptonemal complex protein 3 (Sycp3) gene, exon 5	0.19
4841	AF177242	Bodo saltans clone pBME40 mitocondrial minicircle DNA	0.024
		Homo sapiens elongation of very long chain fatty acids (FEN1/Elo2.	
4842	XM_004347	SUR4/Elo3, yeast)-like 2 (ELOVL2), mRNA	0.66
		Mus musculus phospholipase D1 (PLD1) gene, exons 23 and 24,	
4843	AF083494	complete sequence	0.53
4844	T100/04		
4844	U89684 AB025716	Lycopersicon esculentum protein kinase (LePK7) gene, partial cds	0.22
4043	AB023/16	Staphylococcus aureus gene for FmtB, complete cds	0.57
4846	AF017180	Schizosaccharomyces pombe hexose transporter (Ght2) mRNA, complete cds	
4847	Z26314	P.falciparum gene for STARP antigen	0.87
4848	XM 045160	Homo sapiens KIAA0275 gene product (KIAA0275), mRNA	0.007
10.10	11.12_045100	Mus musculus adult male testis cDNA, RIKEN full-length enriched	11
4849	AK016495	library, clone:4931431C02, full insert sequence	
		Homo sapiens, Similar to hypothetical protein FLJ14058, clone	0.62
4850	BC004496	IMAGE:3831313, mRNA	1.7
4851	XM_005684	Homo sapiens tolloid-like 2 (TLL2), mRNA	1.6
		Sphaerotheca pluvialis cytochrome b (cytb) gene, partial cds,	1.0
4852	AF249077	mitochondrial gene for mitochondrial product	0.047
	XM_016934	Homo sapiens hypothetical protein FLJ12891 (FLJ12891), mRNA	5.4
4854	X65747	R.norvegivus gnat-3 mRNA for gustducin	0.2
4855	XM_011705	Homo sapiens parancoplastic antigen MA2 (PNMA2), mRNA	0.2

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		(Diasav vs. Genbank)	
NO	ACCESSN	DESCRIP	
		Arabidopsis thaliana cytosine-5 methyltransferase mRNA, complete	P VALUE
4856	L10692	cds	
		Arabidopsis thaliana gene for MYB transcription factor Atmyb2,	2.1
4857	AB052250	complete cds, strain: Yo-0	
		Homo sapiens region containing tryptase beta 2; tryptase beta 1	0.36
4859	XM_03353	(LOC82314), mRNA	1.3
		Rattus norvegicus stem cell factor gene, 5' flanking region and exon 1,	1.3
4860	U49846	parnai sequence	0.18
		Tetrahymena thermophila kin2 gene for kinesin-II homologue, exons	0.18
4861	AJ244021	1-4	0.063
			0.005
		Stachyarrhena sp. Jansen-Jacobs 4707 ribosomal protein S16 (rps16)	
4862	AF201021	gene, partial intron sequence; chloroplast gene for chloroplast product	0.057
4863	AJ245636	Mus musculus P2Y1 gene for P2Y1 recentor	0.2
4864	AK023175	Homo sapiens cDNA FLJ13113 fis, clone NT2RP3002590	0.2
		Homo sapiens hypothetical gene supported by AK024355	
4865	XM_034638	(LOC90867), mRNA	1E-08
4866	AB011006	Rattus norvegicus gene for leptin receptor, partial cds	5.5
40.0	*n *	Homo sapiens hypothetical protein DKFZp761I172 (DKFZP761I172),	
4867 4868	XM_031128	MKNA	0.22
4008	AK001442	Homo sapiens cDNA FLJ10580 fis, clone NT2RP2003533	2E-12
4870	AT207054	Saccharomyces servazzii 46.1 kDa protein, KAR4-like protein, and	
4070	AF307954	SPB1-like protein genes, complete cds	0.19
4871	AF055079	Panulirus argus inositol 1,4,5-trisphosphate receptor (IP3R) mRNA, complete cds	
4872	X88849		6
10/2	200047	C.coli ceuB, ceuC, ceuD, ceuE, OrfA, OrfB genes	0.0008
4873	XM 039699	Homo sapiens hypothetical protein FLJ23309 (FLJ23309), mRNA	
		Lactococcus lactis bacteriophage ul36.2 ORF111b (ORF111b),	3E-42
- 1		ORF78b (ORF78b), ORF61b (ORF61b), ORF245 (ORF245), ORF364	ı
- 1		(ORF364), and putative replisome organizer (ORF255) genes,	1
1874	AF212847	complete cds; and ORF241 (ORF241) gene, partial cds	
		(Ora 241) gene, partiai cus	1.9
1875	XM_052223	Homo sapiens DKFZP564G092 protein (DKFZP564G092), mRNA	0.69
1876	AK024606	Homo sapiens cDNA: FLJ20953 ffs. clone ADSF01970	3E-89
		Clavibacter michiganensis subsp. sepedonicus species-specific DNA	JE-67
1877	U96097	clone Cms85	0.47
878	Z48599	L. mexicana Imcpc gene cathepsin B-like cysteine proteinase	0.063
879	NM_014160	Homo sapiens HSPC070 protein (HSPC070) mRNA	5
		Homo sapiens NADH dehydrogenase (ubiquinone) 1 alpha	
880	VIM_031388	subcomplex, 5 (13kD, B13) (NDUFA5), mRNA	0.55
	1 F0 =0 == -	Homo sapiens protein arginine N-methyltransferase 3 (PRMT3)	
881	AF059531	mRNA, partial cds	0.18
882	AF240700	Gymnocrotaphus curvidens cytochrome b (cytb) gene, partial cds,	
552	AL 240 / 00	mitochondrial gene for mitochondrial product	0.65

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	T :		_
		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ II	0	guest (Canada to Combank)	
NO	ACCESSN	DESCRIP	DYVIE
			P VALU
4883	AE005988		2
	1	Neocalanus plumchrus cytochrome oxidase subunit I (CO) gene	+
4884 4885	AF332785	partial cds; mitochondrial gene for mitochondrial product	0.65
4885	AJ278986	Pichia etchellsii plasmid pPE1B	0.055
4886	AL512660	Human DNA sequence from clone RP11-353M9 on chromosome 10,	
1000	AL312000	complete sequence [Homo sapiens]	6.3
4887	AF147082	Homo sapiens gamma-glutamyl hydrolase gene, exons 3 through 7	
		7 armina suprems gamma-gintality) hydrolase gene, exons 3 through 7	2
		Human DNA sequence from clone RP11-563A22 on chromosome 20	
	l	Contains the 5' cnd of a novel gene weakly similar to Xenopus laevis	
4888	AL389874	putative Zic3 binding protein, complete sequence [Homo sapiens]	
4889	AC084157	Caenorhabditis elegans cosmid Y46E12BR, complete sequence	0.008
4891	AE001183	Borrelia burgdorferi (section 69 of 70) of the complete genome	0.084
		genome (occurred to the complete genome	0.66
4892	AE002137	Ureaplasma urealyticum section 38 of 59 of the complete genome	0.50
4893	XM_028606	Homo sapiens Sp1 transcription factor (SP1), mRNA	0.52
4894	NC_001453	Strongylocentrotus purpuratus mitochondrion, complete gamena	6
4895	AK023104	Homo sapiens cDNA FLJ13042 fis, clone NT2RP3001318	0.064
		Homo sapiens stimulated trans-acting factor (50 kDa) (STAF50),	0.064
4896	XM_015243	mRNA (61 aDa) (61 A1 50),	1.7
4897	XM 043823		
4898	AB049900	Homo sapiens hypothetical protein FLJ20958 (FLJ20958), mRNA	0.000004
1070	AD049900	Macaca fascicularis brain cDNA, clone:QnpA-19713	6.1
4899	AE001390	Plasmodium falciparum chromosome 2, section 27 of 73 of the complete sequence	
-027	ALGOISTO	complete sequence	0.23
1900	AE007346	Streptococcus pneumoniae section 29 of 194 of the complete genome	
1901	X17339	Dengue-2 virus NS1 gene for nonstructural protein (patient M2)	0.73
1902	AK000864	Homo sapiens cDNA FLJ10002 ffs, clone HEMBA1000046	0.24
		Panorpa helena cytochrome oxidase I (COI) gene, partial cds;	0.00001
1903	AF180076	mitochondrial gene for mitochondrial product	
		Homo sapiens chromosome 5 clone CTD-2178M23, complete	0.6
904	AC008855	sequence	277.12
905	K02990	Human hepatitis A virus, complete genome	3E-12 0.23
		Escherichia coli O157:H7 EDL933 genome, contig 1 of 3, section 134	0.23
906	AE005310	of 155	0.078
			0.070
907	NM_032717	Homo sapiens hypothetical protein MGC11324 (MGC11324), mRNA	0.68
	1	Homo sapiens, clone MGC:11170 IMAGE:3843148, mRNA	2,30
908	BC004118	complete cds	1.5
	, JI	Human (clone D13-2) L-iditol-2 dehydrogenase gene, exon 4, exon 5,	
909	LEJZJZ I	skon o and exon /	0.0000001
910	0.60117	PLP-A=prolactin-like protein A {5' region, exon 1, intron 1} [rats,	
110	S68117	Genomic, 1187 nt]	0.11

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
4911	NM_016701	Mus musculus nestin (Nes), mRNA	0.21
	1	Lactococcus lactis subsp. lactis IL1403 section 180 of 218 of the	0.21
4912	AE006418	complete genome	0.009
			0.007
4913	XM_041162		4E-50
4914	AF147358	Homo sapiens full length insert cDNA clone YB61E03	0.00003
4915	AE000642	Helicobacter pylori 26695 section 120 of 134 of the complete genome	1.9
4916	XM_040436	Homo sapiens nebulin (NEB), mRNA	2
4917	AB026296	Pisum sativum mRNA for hsr203J homolog, complete cds	0.024
4918	AF159148	Danio rerio Noggin 2 mRNA, complete cds	0.23
		Lucilia cuprina sex-lethal protein SXL2 mRNA, complete cds,	
4919	AF234184	alternatively spliced	0.026
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
4920	AK007229	library, clone:1700121L03, full insert sequence	0.21
4921	AJ404228	Saccharomyces douglasii mitochondrial tRNA gene cluster	0.021
4922	X71604	H.sapiens son-pseudogene	0.53
4000		Homo sapiens DNA (cytosine-5) methyltransferase gene, intron 4,	
4923	AF169122	partial sequence	0.059
4924	70 5 010000	Homo sapiens protein phosphatase, EF hand calcium-binding domain	
4924	XM_040202	1 (PPEF1), mRNA	6E-09
4925	AY027893	Homo sapiens voltage-dependent calcium channel beta 2 subunit	
4923	A102/893	(CACNB2) gene, exon 1	3.3
4926	AF136008	Desulfobacterium sp. BSv41 16S ribosomal RNA gene, partial sequence	
4927	NM 007439	Mus musculus anaplastic lymphoma kinase (Alk), mRNA	5.6
***	1414_007433	with industrius anapiastic lymphoma kinase (Alk), mRNA	0.071
4928	XM 006758	Homo sapiens frizzled (Drosophila) homolog 10 (FZD10), mRNA	
1320	1211_000730	Human DNA sequence from clone RP11-228P1 on chromosome 6,	2
4929	AL139329	complete sequence [Homo sapiens]	
4930	L03188	Saccharomyces cerevisiae integrin analogue gene, complete cds	4E-38
4931	AB039881	Canis familiaris mRNA for cytochrome P450c21, complete cds	0.23
		Mus musculus adult male lung cDNA, RIKEN full-length enriched	0.55
4932	AK004783	library, clone:1200015E14, full insert sequence	6.2
4933	U41343	Human prolargin (PRELP) gene, exon 2	0.0000004
4934	AF157369	Homo sapiens Niemann-Pick C1 protein (NPC1) gene, exon 5	0.000004
4935	XM 016505	Homo sapiens KIAA0999 protein (KIAA0999), mRNA	9E-26
4936	U07807	Human metallothionein IV (MTIV) gene, complete cds	0.008
4937	NM 018818	Mus musculus choroidermia (Chm), mRNA	1.7
		Mus musculus, beta-1,3-N-acetylglucosaminyltransferase 1, clone	
4938	BC009075	MGC:6892 IMAGE:2654354, mRNA, complete cds	1.2
4939		Homo sapiens proteoglycan 3 (PRG3) gene, complete cds	0.24
4940	AK000864	Homo sapiens cDNA FLJ10002 fis, clone HEMBA 1000046	0,00001
		Dictyostelium discoideum developmental protein (DG1105) gene.	
4941	AF076601	partial cds	0.95

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	-	Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		gast (class / to. Combine)	
NO	ACCESSN	DESCRIP	P VALUE
		Bombyx mori insulin receptor-like protein precursor (BIR) mRNA.	FVALUE
4942	AF025542	complete cds	0.64
			0.04
4943	XM_038714	Homo sapiens hypothetical protein dJ37E16.5 (DJ37E16.5), mRNA	6.3
4944	AC016707	Homo sapiens BAC clone RP11-221K4 from Y, complete sequence	
4945	XM 005684	Homo sapiens tolloid-like 2 (TLL2), mRNA	0.054
4946	AJ276170	Rattus norvegicus partial Vps541 processed pseudogene, 3' end	1.1
4947	AJ132986	Vibrio sp. partial 16S ribosomal RNA, isolate A515	0.041 5.9
4948	XM 009513	Homo sapiens hypothetical protein (HSPC207), mRNA	0.21
4949	AF181668	Homo sapiens clone 173p17 polymorphic microsatellite sequence	0.21
-		Dictyostelium discoideum protein kinase YakA (yakA) mRNA,	0.17
4950	AF045453	complete cds	0.007
		Dictyostelium discoideum spore coat structural protein SP65 (cotE)	0.007
4951	AF279135	gene, complete cds	0.003
		Mus musculus 10, 11 days embryo cDNA, RIKEN full-length	0.005
4952	AK012631	enriched library, clone:2810001G20, full insert sequence	0.071
4953	AB047630	Macaca fascicularis brain cDNA, clone:QnpA-13555	0.69
		Hordeum vulgare chloroplast rps15 (partial), ndhH, ndhI, ndhG	
4954	AJ011848	ndhE, psaC, ndhD and ndhA genes	0.054
4955	U34363	Plasmodium falciparum CTRP gene, complete cds	0.18
		Homo sapiens acid ceramidase (ASAH) gene, exons 5 through 14, and	
4956	AF220175	complete cds	0.0009
4958	D26442	Yeast gene for PES4 PAB-like protein, complete cds	0.003
4959	NC_002184	Penaeus monodon mitochondrion, complete genome	1.9
4960	XM_039251	Homo sapiens 37381 (KIAA1660), mRNA	5
1001	70 ( 004460	Homo sapiens signal sequence receptor, alpha (translocon-associated	
4961 4962	XM_004462	protein alpha) (SSR1), mRNA	0.72
4902	AF252420	Bos taurus clone MNB-5 microsatellite sequence	0.08
4963	XM_035186	Homo sapiens RCE1 (S. Cerevisiae) homolog, prenyl protein protease	
4903	VINT_022190	(RCE1), mRNA	2
4964	AL110134	Homo sapiens mRNA; cDNA DKFZp564H072 (from clone DKFZp564H072)	
4204			0.058
4965	AL049452	Homo sapiens mRNA; cDNA DKFZp586C1322 (from clone DKFZp586C1322)	
4966		Macaca fascicularis brain cDNA, clone:QnpA-12025	0.67
4967	XM 043315	Homo sapiens KIAA0229 protein (KIAA0229), mRNA	0.077
-		Mus musculus 8 days embryo cDNA, RIKEN full-length enriched	6.2
4968	AK017495	library, clone:5730405K23, full insert sequence	0.025
		Candida glabrata TEF3 gene for translation elongation factor3,	0.023
4969	AB012141	complete cds	0.003
4970	AK022914	Homo sapiens cDNA FLJ12852 fis, clone NT2RP2003445	0.0003
		Homo sapiens COT kinase proto-oncogene, partial cds, alternatively	
4971	AF133211	spliced	0.008

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	T		
		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		Jan Sandas)	
NO	ACCESSN	DESCRIP	P VALUE
		Simian immunodeficiency virus strain SIVrcmNg409 from Nigeria 5'	FVALUE
1	ł	long terminal repeat, partial sequence: gag protein (gag) gang	
		complete cds, pol protein (pol) gene, partial cds, and vif gene,	
4972	AF349681	complete sequence	0.68
1		Homo sapiens fibrinogen gamma chain precursor (FGG) gene,	0.08
4973	AF350254	complete cds, alternatively spliced	0.21
4974	AF314052	Rhizobium sp. CIAT613 nifHc gene, partial sequence	1.9
		Homo sapiens inhibitor of DNA binding 4, dominant negative helix-	
4975	XM_036277	loop-helix protein (ID4), mRNA	0.076
	l	Homo sapicns 17-beta-hydroxysteroid dehydrogenase IV (HSD17B4)	
4976	AF057740	gene, exon 24 and complete cds	0.00003
4000	7D 4 4	Homo sapiens similar to NAD(P) dependent steroid dehydrogenase-	
4977	XM_051777		4E-11
4978	AF314052	Rhizobium sp. CIAT613 nifHc gene, partial sequence	2
4979	DC010011	Homo sapiens, Similar to adaptor protein containing pH domain, PTB	
49/9	BC010911	domain and leucine zipper motif, clone IMAGE:4295177, mRNA	0.076
4980	AF214529	T	
4981	AK024235	Homo sapiens dystrophin (DMD) gene, intron 51, partial sequence	0.001
7701	AIX024255	Homo sapiens cDNA FLJ14173 fis, clone NT2RP2002755	e-104
4982	AB035089	Homo sapiens SCCA2 gene for squamous cell carcinoma antigen 2, complete cds	
4983	M97555	Schistosoma mansoni tropomyosin mRNA, complete cds	0.23
	11277333	Drosophila melanogaster genomic scaffold 142000013385404,	0.22
4984	AE002745	complete sequence	
4985	X99288	B.calamita repeat region, clone-lib micro-8	0.003
4987	AJ308840	Coffea arabica microsatellite DNA, clone 16-6CTG	0.23
		Cucurbita moschata CmATS1;1 gene for acyl-(acyl-carrier-protein):	0.025
4988	AB049134	glycerol-3-phosphate acyltransferase, complete cds	0.53
4989	X03991	Human glucagon gene	0.53
		Arabidopsis thaliana zinc finger-like protein (F3C22_200) mRNA,	0.074
4990	AY042871	complete cds	0.077
			0.077
4991	AF214529	Homo sapiens dystrophin (DMD) gene, intron 51, partial sequence	0.001
		Staphylococcus epidermidis strain SR1 clone step.1002a02 genomic	
4992	AF269319	sequence	2
4993	AB009801	Homo sapiens gene for osteonidogen, intron 2	0.69
		Torulopsis glabrata metal-activated transcription factor (amt1) gene,	
4994	M69146	complete cds	0.65
1005		Human DNA sequence from clone 346P11 on chromosome Xq21.2-	
4995	AL121822	21.33, complete sequence [Homo sapiens]	5E-09
1006	A.Formore	Lactuca sativa resistance protein candidate RGC2N (RGC2N)	
4996 4997	AF072273	pseudogene, complete sequence	0.22
4397	AL358273	S.pombe chromosome II cosmid c29C10	0.22
4998	BC010011	Homo sapiens, Similar to RIKEN cDNA 4931428D14 gene, clone	
7/70	DC010011	MGC:15407 IMAGE:4309613, mRNA, complete cds	0.22

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRI	P VALUE
4999	AF090889	Homo sapiens clone HQ0092	0.71
5000	M93140	Glycine max cv Prize protein kinase mRNA	2.1
5001	AE005266	Escherichia coli O157:H7 EDL933 genome, contig 1 of 3, section 90 of 155	1.8
5002	AE007350	Streptococcus pneumoniae section 33 of 194 of the complete genome	0.23
<b>5</b> 003	BC008973	Homo sapiens, transmembrane protein with EGF-like and two follistatin-like domains 2, clone MGC:17007 IMAGE:4180709, mRNA, complete cds	0.023
5004	X13439	Cochliobolus heterostrophus mitochondrial gene for ATPase subunit 6	0.19
		HIV-1 p5c028-h from USA envelope glycoprotein (env) gene, partial	9122
5005	AF138069	cds	2.1
5006	NM_002847	Homo sapiens protein tyrosine phosphatase, receptor type, N polypeptide 2 (PTPRN2), mRNA	6
		Homo sapiens hypothetical gene supported by X04201; X04588;	
		AK026559; AY004867; BC000771; BC008407; BC008425	
5007	XM_036827	(LOC91191), mRNA	1
5008	NM_031389		1.7
5009	BC010355	Homo sapiens, Similar to nuclear protein, clone IMAGE:4248516, mRNA	5.4
5010	AE003909	Xylella fastidiosa 9a5c, section 55 of 229 of the complete genome	5.1
5011	NM_031389	Mus musculus ribonuclease/angiogenin inhibitor 2 (Rnh2), mRNA	1.7
5012	XM 016747	Homo sapiens latent transforming growth factor beta binding protein 1 (LTBP1), mRNA	
5012	2111_010747	Schistosoma mansoni homeodomain protein (smox-5) mRNA,	2E-10
5013	M85304	complete cds	0.009
			0.009
5014	NC_001398	Saccharomyces cerevisiae 2 micron circle plasmid, complete sequence	0.65
		Drosophila melanogaster rad21 mitotic cohesin (drad21) gene, exons	0.05
5015	AF267746	3 through 8 and complete cds	0.003
	XM_052430	Homo sapiens E3 ubiquitin ligase SMURF1 (SMURF1), mRNA	2E-30
5017	X67813	C.familiaris SRP72 mRNA for signal recognition particle	0.16
5018	AF031848	Dermacentor marginatus 12S ribosomal RNA gene, mitochondrial gene for mitochondrial RNA, partial sequence	0.061
5019	XM_003317	Homo sapiens PDZ domain containing guanine nucleotide exchange factor(GEF)1 (PDZ-GEF1), mRNA	0.67
5020	XM 030673	Homo sapiens methyl-CpG binding domain protein 4 (MBD4), mRNA	0.6
	050075	Saccharomyces cerevisiae nuclear-encoded mitochondrial isoleucyl-	0.0
5021	L38957	tRNA synthetase (ISM1) gene, complete cds	2.1
		Hepatitis C virus isolate SCpreSC2c14 nonfunctional polyprotein	
5022	AF345089	gene, partial sequence	0.59

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-			Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEC			(Blash Vs. Genbank)	
N		ACCES	SN DESCRIP	
50:	23	AC0911	24   Caenorhabditis elegans cognid V4747	P VALI
1			E.octocarinatus RPA2 gene for RNA polymerase I second largest	1.7
502	24	X6645	subunit subunit	
	$\neg$		Homo saniens hypothetical	0.066
_ 502	5	XM_0105	Homo sapiens hypothetical gene supported by AB033071; AB05148 663 AK000726; NM_015383 (LOC92396), mRNA	0;
502	6	AF25672		1E-15
502	7	AF38560	Homo sapiens minisatellite sequence	0.5
			Pleurotus ostrastva lineau	5.3
502	8	AF35510	Pleurotus ostreatus linear mitochondrial plasmid mlp2, partial sequence	
	$\top$			0.18
5029	9	XM_0348	Homo sapiens similar to hypothetical protein FLJ22294 (H. sapiens) 33 (LOC90918), mRNA	
	+		(LOC50518), IIIRNA	1.8
5030	,	AF37400	S Pottus	1.0
	+	111 37400		8.3
				0,5
5031		AV01444	omoryome ussue CDNA, RIKEN full-length engles at 12	ł
5051	+	AK01444	clone:3830421F13, full insert sequence	0.18
				0.18
5032	1		Human DNA sequence from clone RP11-299G2 on chromosome 6	1
3032	+	AL355577		0.0000
***	1		ivius musculus Chetk-alpha gene for choling/other-land	0.0003
5033	+-	AB030616		
	1		pol Isimian Mason-Pfizer D-type patronics Ch ( pro-	6.1
5034	4	S66671		
•	1		Couldin incasii chloroplast the gone for situal	0.008
5035	1	AB038481		
	1		Rattus norvegicus Sprague-Dawley heme oxygenase-2 non-reducing	0.22
5036	L	U05013		
			Homo sapiens (subclone 1_h7 from P1 H69) DNA sequence, complete sequence	1.9
5037		L81904	sequence sequence, complete	
				1.7
5038	A	F242384	Homo sapiens phytanovil Co. t. t.	
5039		X52863	Homo sapiens phytanoyl-CoA hydroxylase (PHYH) gene, exon 7 G.max Gy4 gene for glycinin	1.9
			Homo serviene endersi	5.9
040	X	4_002625	Homo sapiens ankyrin repeat and SOCS box-containing 3 (ASB3), mRNA	
041		J00789		6
$\neg$	_		Mus musculus Dp71 (DMD) gene, partial cds	0.068
042	Al	3017429	Oryza sativa gene for mitochondrial ribosomal portein S14, succinate	
			dehydrogenase iron-protein subunit (SDHB)	2
043	т	67518	Mathanagaa	
044		048782	Methanococcus jannaschii section 60 of 150 of the complete genome	0.002
045		3055263		0.2
	- 7.1	033203	Macaca fascicularis brain cDNA, clone:QfIA-10674	0.003
046		65281		0.003
-~-			drebrin {clone eDcg5} [chickens, Genomic, 160 nt, segment 3 of 10]	0.61
47	4.7			10.0
4/	AF			
48		06266	Anthirrhinum majus DNA for transposon tam2	0.25

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- '		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			T
NO	ACCESSN	DESCRIP	P VALU
5049	AF272387	Homo sprions leveles -i	
5050	NC 00158		7E-22
		resistant populoinavitus type 34, complete genome	0.25
5051	NM_01707	Rattus norvegicus Acetylcholine receptor alpha 5 (Chrna5), mRNA	0.00
			0.64
5052	XM_030510		2
5053	Z48433	L.odomensis vicilin pseudogene	0.047
5054	3.631004	P.hybrida 5-enolpyruvylshikimate 3-phosphate synthase mRNA,	
3034	M21084	complete cds	0.2
5055	U67518	Methanococcus in monetiii' co case	
	00.010	Methanococcus jannaschii section 60 of 150 of the complete genome	0.002
5056	AF215555	Blakea trinervia NADH dehydrogenase subunit F (ndhF) gene, partial cds; chloroplast gene for chloroplast product	
		Plasmodium falciparum chromosome 2, section 16 of 73 of the	0.63
5057	AE001379	complete sequence	5.4
.		Candida albicans heat shock protein Hsp104 (HSP104) gene, HSP104-	3.4
5059	AF362391	Ib allele, complete cds	4.4
5060	AJ000056	Homo sapiens tyrosine aminotransferase gene, 5' flanking region	0.57
5061	A77110541	Fil V-1 Isolate 08108v3 from USA, envelope glycoprotein (env) gene	
1001	AF112541	partial cds	0.007
- 1		District	
		Diadasia consociata cytochrome oxidase subunit I gene, partial cds;	
5062	AF300534	tRNA-Leu gene, complete sequence; and cytochrome oxidase subunit II gene, partial cds; mitochondrial genes for mitochondrial products	
5063	J00571	Mouse Ig kappa unproductively rearranged V-T2; V-J region	0.22
5064	X95275	P.falciparum complete gene map of plastid-like DNA (IR-A)	0.001
5065	Y09746	H.oligactis mRNA for heat shock protein 70	0.062
066	X95536	H.sapiens earl gene	0.46
1		C.ellipsoidea rbcL, rps14, trnM, trnG, trnD, trnS, rps4, atpE, atpB	2E-64
067	D10997	genes, complete cds	1.8
5068	Z73968	Caenorhabditis elegans cosmid C05A2, complete sequence	0.67
	XM_012733	Homo sapiens KIAA1012 protein (KIAA1012) mPNIA	0.24
070	AE000745	Aquifex aeolicus section 77 of 109 of the complete genome	0,21
071		Diploptera punctata high-affinity Na+-dependent alutamate	
0/1	AF208521	transporter (EAAT1) mRNA, complete cds	0.08
072 3	CM 032656	Homo sapiens mannosidase, alpha, class 2B, member 1 (MAN2B1), mRNA	
073		S.cerevisiae chromosome XI reading frame ORF YKL063c	1.8
074 3	CM 050445	Homo sapiens KIAA0164 gene product (KIAA0164), mRNA	6,2
		suprementation gene product (KIAA0164), mRNA	0.66
	ŀ	Pectobacterium chrysanthemi ArgG (argG) gene, partial cds;	1
	1	regulatory protein PecS (pecS), regulatory protein PecM (pecM),	j
	li li	ndigoidine systhesis protein IdgA (idgA), and indigoidine systhesis	
75	AF265211	protein IdgB (idgB) genes, complete cds; and indigoidine systhesis	
		and indigordine sys	0.72

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1.	1		
		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ I			
NO	ACCESSN		P VALU
5000	1	Borrelia burgdorferi plasmid, orfA, B, C, D, F, bdrV genes, clone	I VALU
5076	X87201	IPONIB14 and POMB17	0.076
	1	Mus musculus myelin-associated oligodendrocytic basic proteins	0.076
5077		[MODE 170 and MOBPO9 (Mohn) gene evone 2 through 5 1	1
3077	AF120475	complete cus, afternatively spliced	0.0009
5078	15050105	Staphylococcus epidermidis strain SR1 clone step.4050g06 genomic	0.0005
3078	AF270407	sequence	0.67
5079	U47138	Inkoo virus SW AR 83-161 nucleocapsid protein and non-structural	
3019	04/138	iprotein genes, complete eds	0.16
5080	AB049420	Mus musculus DNA, clone:7-1, derived from 4x chromosome of	
2000	AD049420	T(X,4)37H translocation	6.5
	l	Chair I	
	!	Staphylococcus xylosus choline transporter (cudT), putative regulatory	
5081	AF009415	protein (cuic), grycine betaine aldehyde dehydrogeness (cuid )	ł
5001	71 005415	chounic denydrogenase (CudB) genes complete ede	0.075
5082	AE001395	Plasmodium falciparum chromosome 2, section 32 of 73 of the complete sequence	
	1112001393	Describite and	0.24
5083	U62542	Drosophila melanogaster dead ringer (dead ringer) mRNA, complete cds	
	- 002012		0.12
5084	AL445527	Human DNA sequence from clone RP11-115C2 on chromosome 1, complete sequence [Homo sapiens]	
5085	XM 016052	Homo sapiens LOC86766 (LOC86766), mRNA	5.5
		Homo sapiens hypothetical gene supported by J04178 (LOC91343),	0.24
5086	XM 037781	ImRNA (LOC91343),	
5087	X96932	N.tabacum gene encoding ascorbate oxidase-related protein	6.4
5088	AJ006986	Streptococcus pneumoniae type 33F DNA, capsular gene cluster	1.6
5089	M60858	Human nucleolin gene, complete cds	0.019
5090	AK026299	Homo sapiens cDNA: FLJ22646 fis, clone HSI07178	0.66
		Homo sapiens PAC clone RP5-1188N21 from 7q11.23-q21.1,	0.21
5091	AC006025	complete sequence	
5092	X03100	Human HLA-SB(DP) alpha gene	0.67
5093	AF181671	Homo sapiens clone 77i13 polymorphic microcatellite some	1.8
T		Dictyostelium discoideum filament-interacting protein (fip) mRNA,	0.055
094	111 330000	complete cus	0.10
i		Human Y-chromosome RNA recognition motif protein (YRRM) gene,	0.19
095	030019	exon 5, partial cds, subclone 782	0.023
		Homo sapiens intrinsic factor-vitamin B12 receptor (CUBN) gene,	0,023
096	1M 243111	exous 32 and 53	2
097	XM_050127	Homo sapiens oligophrenin 1 (OPHN1), mRNA	0.5
- 1	-	Drosophila melanogaster mRNA for LIM domain protein (espinas	0.5
098	14251652	gene)	0.072
199	AF036696	Caenorhabditis elegans cosmid F15B10	0.072
	[]	Mus musculus phospholipase D2 gene, exons 13 through 25 and	0.2
100	111 032234	ompiete cas	5.7
101	AK026741 I	Iomo sapiens cDNA: FLJ23088 fis, clone LNG07026	0.003
102	K02212 I	duman alpha-1-antitrypsin gene (S variant), complete cds	1.7

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ II	O	Treatest Prognoti (Biastiv vs. Genoank)	
NO	ACCESSN	DESCRIP	
5103	AB055370	Macaca fascicularis brain cDNA, clone:QflA-12708	P VALU
		Mus musculus adult male stomach cDNA, RIKEN full-length	0.079
5104	AK008762	enriched library, clone:2210021122, full insert sequence	1
5105	M71245	Rat prostatein C3 subunit gene, complete cds	2.1
5106	AY007089	Homo sapiens clone TCCCIA00110 mRNA sequence	0.69
5107	AF113007	Homo sapiens PRO0066 mRNA, complete eds	0.000009
5108	AF125461	Caenorhabditis elegans cosmid Y8A9A, complete sequence	0.0007
		Rattus norvegicus putative pheromone receptor (Go-VN2) mRNA,	0.69
5109	AF016179	complete cds	1
5110	AK021604	Homo sapiens cDNA FLJ11542 fis, clone HEMBA1002801	0.071
5111	Z73968	Caenorhabditis elegans cosmid C05A2, complete sequence	0.026
5112	U40995	Danio rerio Hox-a1 gene, complete cds	0.54
5113	NC 001728	Odontoglossum ringspot virus, complete genome	0.009
_		Guornogiossum ringspot virtis, complete genome	0.027
			gi 8778469
			gb AAF794
		i	77.1IAC02
- 1		,	2492_21
1			(AC022492
İ	2055 740		) FIL3.30
114	2056.P10.gz4		[Arabidops
115	3_268157 AF234932	8778469	is thaliana]
116	M60858	Chtenopteryx sicula clone 9 actin gene, partial cds	0.24
110	100858	Human nucleolin gene, complete cds	0.66
117	XM_003100	Homo sapiens hypothetical protein FLJ11142 (FLJ11142), mRNA	0.23
118	AF129278	Buchnera aphidicola natural-host Diuraphis novia gluconate-6-	
119		phosphate dehydrogenase (gnd) gene, partial cds	6.4
120	AF283665 AJ005205	Heterodera glycines guanylyl cyclase mRNA, complete cds	5.4
121	AF135472	Homo sapiens 5HT3 gene for serotonin 3 receptor	0.24
122	XM 036007	Clostridium beijerinckii Rep protein gene, complete cds	0.23
122	AM_036007	Homo sapiens transmembrane protein 1 (TMEM1), mRNA	2.1
123	AL031426	Human DNA sequence from clone CTA-191D12 on chromosome 22q13.1 Contains two exons of the APOL2 gene for apolipoprotein L 2, ESTs and GSSs, complete sequence [Homo sapiens]	0.000001
	XM_049821	Homo sapiens hypothetical protein MGC14832 (MGC14832) mPNA	5.4
	AM_028344	Homo sapiens ribosomal protein L5 (RPL5) mRNA	0.16
26	AF016653	Caenorhabditis elegans cosmid C41D7, complete sequence	0.16
Т	,	H. sapiens telomeric DNA sequence, clone 8QTEL025, read	0.7
127	Z96752	8QTELOO025.seq	0.006
128		Sus scrofa (pig) apomucin mRNA, 3' end	0.006
		a a -pointent milato, 5 cilu	0.026

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	1 .		
		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ II		rearest reignoof (Blastin vs. Genbank)	
NO	ACCESS		D ****
		Pentachondra pumila specimen-voncher Cherry 96/1 metures (m. 47)	P VALU
5129	AY00510	Igene, partial cds; chloroplast gene for chloroplast product	
1		Homo sapiens beaded filament component protein (CP49) gang	2.2
5130	AF195044	I partial cus	0.57
5131	AK021963		0.0005
5132	AF034085	Caenornabditis elegans UNC-45 (unc-45) gene complete ed-	0.0001
5133	******	Mus musculus L1 neural cell adhesion protein gene 5' sequence and	
5134	U91929	EXOII 1	2.1
3134	NM_01078		0.68
5135	ABOLTOLO	Mus musculus gene for gross cystic disease fluid protein 15, complete	
3133	AB017918	cas	0.001
5136	U55243	Dictyostelium discoideum glyceraldehyde-3-phosphate dehydrogenasc	
5137	AF061296	(CALDA) gene, partial cos	0.24
3137	A1'001296	Homo sapiens sulfonylurea receptor (SUR2) gene, exon 9	9E-45
		Liuman Dhia C	
5138	AL050314	Human DNA sequence from clone 100G10 on chromosome 22q13.31-	
	142000314	13.33. Contains GSSs, complete sequence [Homo sapiens]	6.4
5139	XM 036394	Homo sapiens calcium/calmodulin-dependent protein kinase IV (CAMK4), mRNA	
5140	XM 047419		0.003
5141	AJ404314	Amphisorus hemprichii 18S rRNA gene, isolate 720	0.21
		Homo sapiens myosin phosphatase target subunit 2 (MYPT2) gene,	0.71
5142	AF324889	exon 1	
		Bacteriophage PBS2 (from B.subtilis) uracil-DNA glycosylase	0.23
5143	J04434	Inhibitor gene, complete eds	0.61
i		Homo sapiens inter-alpha (globulin) inhibitor, H1 polypeptide	0.61
5144	XM_041213	(ITIH1), mRNA	5,5
			5,5
5145	AF187881	Homo sapiens leukointegrin alpha d gene, promoter and partial cds	0.026
5146 5147	Z70180	P.quinquangularis chloroplast matK gene	2
3147	AK025364	Homo sapiens cDNA: FLJ21711 fis, clone COL10156	0
5148	V3 6 045504	**	
3148	XM_045594	Homo sapiens hypothetical protein MGC4816 (MGC4816), mRNA	5E-82
5150	AB029068		
3130	AD029068	Mus musculus gene for Ser/Thr kinase KKIAMRE, exon 1, 2, 3, 4, 5	0.63
5151	AF215555	Diakea triffervia NADH dehydrogenase subupit E (ndbE) gone martial	
	111 2 1 3 3 3 3	cds; chloroplast gene for chloroplast product	0.71
5152	XM 026967	Homo saniens hypothetical	
5153	M32328	Homo sapiens hypothetical protein MGC2668 (MGC2668), mRNA C.pipiens esterase B1 gene, complete cds	1.9
154	U58744	Caenorhabditis elegans cosmid F55F1	0.008
_		(Sen'll) AgaR (agaR), KbaZ (kbaZ), AgaV (agaV), AgaW	0.43
- 1		(agaW), AgaE (agaE), AgaF (agaF), AgaA (agaA), AgaS (agaS),	1
		(kbaY), AgaB (agaB), AgaC (agaC), AgaD (agaD), and AgaI	1
155	AF228498	(agal) genes, complete eds	0.72
			0.72

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
5156	L36898	Saccharomyces cerevisiae mitochondrion transfer RNA-Glu (tRNA-Glu) gene	0.69
5158	D88262	Pisum sativum PsCHS5 gene for chalcone synthase, complete cds	0.7
5159	AK018080	Mus musculus 11 days embryo head cDNA, RIKEN full-length enriched library, clone:6230403H02, full insert sequence	
5160	X95275	P.falciparum complete gene map of plastid-like DNA (IR-A)	6.4
5161	D16417	Dictyostelium discoideum mRNA	0.022
5162	AF267204	Candidatus Carsonella ruddii natural-host Cacopsylla brunneipennis ATP synthase alpha subunit (atpA) gene, partial cds, ATP synthase gamma subunit (atpG) gene, complete cds; and ATP synthase beta subunit (atpD) gene, partial cds	
5163	AF264650	Asterella tenella trnL gene, intron sequence; chloroplast gene for chloroplast product	0.026
5164	AF003143	Caenorhabditis elegans cosmid C53H9, complete sequence	0.23
5165	M19459	Plasmodium falciparum interspersed repetitive DNA and an open reading frame	0.008
5166	AK001362	Homo sapiens cDNA FLJ10500 fis, clone NT2RP2000369	0.008
5167	AF170023	Anopheles gambiae clone I38 genomic sequence	0.57
5168	AJ012333	Buchnera aphidicola plasmid pBTc2, trpE and trpG genes	0.006
5169	AF158077	Pisacha naga 16S ribosomal RNA gene, partial sequence; mitochondrial gene for mitochondrial product	0.019
5170	U68721	Botryotinia fuckeliana endopolygalacturonase 5 (BcPGA5) gene, complete cds	0.21
5171	AK026372	Homo sapiens cDNA: FLJ22719 fis, clone HSI14307	e-116
5172	NM_031130	Rattus norvegicus nuclear receptor subfamily 2, group F, member 1 (Nr2f1), mRNA	3
5173	AY034470	Homo sapiens clone BGL2 mRNA sequence	0.064
5174	AF108841	Homo sapiens human endogenous retrovirus HERV-H10 pol protein (pol) gene, partial cds; env pseudogene and 3' LTR, complete sequence	0,009
5175	AJ251957	Ciona intestinalis mRNA for nuclear lamin (lamin L2 gene)	1.9
5176	XM_045437	Homo sapiens a disintegrin and metalloproteinase domain 33 (ADAM33), mRNA	0.71
5177	AF177983	Homo sapieus NAD+-dependent 15-hydroxyprostaglandin dehydrogenase (PGDH) gene, promoter, exons 1 and 2 and partial cds	9.9
5178	AF181632	Drosophila melanogaster BcDNA.GH05095 (BcDNA.GH05095) mRNA, complete cds	0.23
5179		Mus musculus 10 days neonate cerebellum cDNA, RIKEN full-length euriched library, clone:6530414F19, full insert sequence	2.1
5180	XM_050838	Homo sapiens wingless-type MMTV integration site family, member 10A (WNT10A), mRNA	0.22
5181	AJ308588	Medicago truncatula partial mRNA for nodulin 25 (nod25 gene), clone 3	4.3

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID	- I		
NO	ACCESSN	DESCRIP	P VALU
		Homo sapiens cis-golgi SNARE (GOSR1) gene, exon 9 and partial	F VALU
5182	AF229797	cds	1 0
5185	Z68107	Caenorhabditis elegans cosmid F48C5, complete sequence	2.1
		Mus musculus germ cell-specific unknown protein mRNA, complete	2.1
5186	AY026045	cds	0.079
			0.075
5187	AF309515	Staphylococcus aurcus serine protease operon, complete sequence	1
		Pleurotus ostreatus linear mitochondrial plasmid mlp2, partial	
5188	AF355103	sequence	0.069
	1	Panulirus argus inositol 1,4,5-trisphosphate receptor (IP3R) mRNA,	
5189	AF055079	complete cds	5.2
		Arabidopsis thaliana chromosome II section 163 of 255 of the	
5190	AC007184	complete sequence. Sequence from clones T11P11	6.1
5191	AB044348	Arabidopsis thaliana AtSUG1 mRNA, complete cds	1.2
5192	XM_003320	Homo sapiens hypothetical protein FLJ11155 (FLJ11155), mRNA	5.3
		Legionella fallonii macrophage infectivity potentiator (mip) gene,	
5193	AF148987	partial cds	0.067
5194	X54200	Chicken mRNA for GARS-AIRS-GART	0.019
5195			
5195	AE005885	Caulobacter crescentus section 211 of 359 of the complete genome	0.62
i		Escherichia coli AgaR (agaR), KbaZ (kbaZ), AgaV (agaV), AgaW	
		(agaW), AgaE (agaE), AgaF (agaF), AgaA (agaA), AgaS (agaS),	
5196	AF228498	KbaY (kbaY), AgaB (agaB), AgaC (agaC), AgaD (agaD), and AgaI	
5197	U09859	(agaI) genes, complete cds	0.63
3197	009839	Bos taurus enterokinase mRNA, complete cds	6.2
5198	AF057708	Populus balsamifera subsp. trichocarpa PTD protein (PTD) gene, complete cds	
5199	AF238381	Homo sapiens PTOV1 (PTOV1) gene, complete cds	0.68
24,7,7	711 250501	Plasmodium falciparum para-aminobenzoic acid synthetase gene,	0.18
5200	AF119554	complete cds	
5201	X82675	B. taurus BoLA-A11 gene (exon 6-8)	0.13
		D. Maria Dolla-ATT gene (exoli 0-8)	0.71
5202	AF303087	Tilapia mossambica glycoprotein alpha subunit mRNA, partial cds	0.64
		Homo sapiens serologically defined colon cancer antigen 16	0.64
5203	NM_006649	(SDCCAG16), mRNA	1.7
5204	AF240597	Homo sapiens clone 7ptelc11t7 sequence	e-112
		Mus musculus, RIKEN cDNA 4930529O08 gene, clone MGC:7935	6-112
5205	BC006702	IMAGE:3583936, mRNA, complete cds	6.3
5206	AF295769	Mus musculus SMAD1 gene, intron 1, partial sequence	0.007
		Toxoplasma gondii PITSLRE-like protein kinase (tpk1) gene, partial	0.007
5207	AF118099	cds	1.9
5208	AF143952	Homo sapiens PELOTA (PELOTA) gene, complete cds	0.44
		Influenza A virus (A/HongKong/156/97(H5N1)) nucleoprotein	V. 1-7
5209	AF028710	mRNA, complete cds	1.7
5210	XM 017097	Homo sapiens active BCR-related gene (ABR), mRNA	0.7

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		Jan ( Sandalat)	
NO	ACCESSN	DESCRIP	P VALUE
5211	M33312	Rat hepatic steroid hydroxylase IIA1 (CYP2A1) gene, complete cds	0.54
5212	L05616	Dictyostelium purpureum (Dpp4) DNA sequence, repeat region	0.061
		Plasmodium falciparum chromosome 2, section 62 of 73 of the	0.001
5213	AE001425	complete sequence	0.001
		Homo sapiens similar to plakophilin 2 (H. sapiens) (LOC93271),	
5214	XM_006574		0.009
5215	XM_011195	Homo sapiens centromere protein E (312kD) (CENPE), mRNA	0.002
5216	AF181881	Helicobacter wileyi T-ID (4-ID) - 1 DAY	
0210	111 101001	Helicobacter pylori TolB (tolB) and PAL (excC) genes, complete cds Human B-lymphocyte activation antigen (B7.1) gene, enhancer region	0.65
5217	U33208	and 5' UTR	
5218	AB052187	Macaca fascicularis brain cDNA, clone:QnpA-12170	0.0003
		Drosophila melanogaster H.M.S. Beagle transposon long terminal	0.7
		repeat, complete sequence; and heat shock protein Hsp70Ab gene,	
5219	AY032741	promoter and partial cds	
		Pisum sativum lip1 mutant cop1 gene for constitutively	0.63
5220	AJ276592	photomorphogenic I protein, exons 1-19	
$\neg \neg$		Homo sapiens similar to cadherin related 23 (H. sapiens)	5,7
5221	XM_045136	(LOC92444), mRNA	2E-52
		Mus musculus 0 day neonate skin cDNA, RIKEN full-length enriched	26-32
5222	AK014534	library, clone:4631424J17, full insert sequence	2E-55
5223	U67572	Methanococcus jannaschii section 114 of 150 of the complete genome	
		Homo sapiens cDNA: FLJ22954 fis, clone KAT09813, highly similar	0.65
5224	AK026607	to AF010315 Homo sapiens Pig11 (PIG11) mRNA	0.21
5225	Y08580	F.rubripes hsp70-3 gene, 3'UTR	1.8
		Human DNA sequence from clone RP11-109B17 on chromosome 1.	1.8
5226	AL359819	complete sequence [Homo sapiens]	0.76
		Toxoplasma gondii PITSLRE-like protein kinase (tpk1) gene, partial	0.70
5227	AF118099	cds (4111) golds platen	1.9
		Homo sapiens amiloride-sensitive epithelial sodium channel gamma	
5228	U48937	subunit gene, promoter region and exon 1	5.1
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
5229	AK017057	library, clone:4933434120, full insert sequence	1.8
5230	AF044770	Phaeoprogne tapera microsatellite HrU6 allele 2 repeat region	4.6
5231	XM_012268	Homo sapiens amino acid transporter 2 (KIAA1382), mRNA	0.009
5232	BC008544	Mus musculus, Similar to hypothetical protein, clone MGC:7703 IMAGE:3497634, mRNA, complete cds	
		Sapporo-like virus Yak2-2000-jp pol gene for RNA-dependent RNA	7E-22
5233	AB046353	sapporo-like virus Yakz-2000-jp poi gene for RNA-dependent RNA polymerase, partial cds	6.2
			0.2
5234	L07305	Histoplasma capsulatum (clone pMS3) H-ATPase gene, complete eds	0.65
5235 1	NM 012828	Rattus norvegicus Calcium channel subunit beta 3 (Cacnb3), mRNA	2
5236		Helicobacter pylori strain 96-74 BabB (babB) gene, partial cds	0.71
		F7-5.7 Strain 20-74 Datio (Gauss) gene, partial cds	0.71

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	T		<del></del>
		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		H. sapiens (D8S266) DNA segment containing (CA) repeat; clone	1 11202
5237	Z16681	AFM151ye3; single read	0.003
		Staphylococcus epidermidis composite transposon DNA, complete	
5238	AB049452	sequence	0.077
5239	X83673	X.laevis SSB1 gene	1.5
		Homo sapiens serine (or cysteine) proteinase inhibitor, clade F (alpha-	
5240	373 6 0 4 5 1 0 0	2 antiplasmin, pigment epithelium derived factor), member 2	
5240	XM_045182		6.2
5241	AF263074	Cobitis paludica cytochrome b gene, partial cds; mitochondrial gene	
5242	XM_029785	for mitochondrial product	0.15
3242	AM_029/85		2.1
5243	AE006148	Pasteurella multocida PM70 section 115 of 204 of the complete genome	1
5244	X81975		5.5
5245	NM 033105	B. vulgaris mRNA for sucrose 6-phosphate synthase	1.3
3243	14M_033103	Homo sapiens beta cysteine string protein (LOC85479), mRNA	0.000006
5246	AC006164	Home coniens class ITHICC. 20	
5247	AJ279971	Homo sapiens clone UWGC:y28gap from 6p21, complete sequence Canis familiaris microsatellite DNA, clone DTRcn.20	0.47
3247	Pazissii	Escherichia coli O157:H7 EDL933 genome, contig 3 of 3, section 237	0.022
5248	AE005618	of 290	
52.0	1112003018	Listeria monocytogenes NotI restriction site number two and flanking	0.56
5249	AJ012338	Isequences	1.6
5251	D14027	Mouse DNA for TL antigen, complete cds	5
5252	Y12285	H.armigera mRNA for putative serine protease, clone SR106	0.009
5253	AF281912	Homo sapiens clone 16qtel_c89bt7 sequence	1E-68
5254	NC_001790	Ovine papillomavirus 2, complete genome	0.61
5255	XM_031110	Homo sapiens kallikrein 10 (KLK10), mRNA	0.041
		Homo sapiens guanine nucleotide binding protein (G protein), alpha	
5256	XM_042732	inhibiting activity polypeptide 3 (GNAI3), mRNA	0.42
5257	XM_033541	Homo sapiens KIAA0704 protein (ORP3), mRNA	0.022
1		Human DNA sequence from clone RP11-245H22 on chromosome 6	
5258	AL512432	Contains GSSs, complete sequence [Homo sapiens]	0.0001
		Human adenovirus type 9 E4 protein (E4), Orf2, Orf3, Orf4, and Orf6	
5259	S82508	genes, complete cds; and Orf7 gene, partial cds	1.6
5260	XM_015579	Homo sapiens hypothetical protein FLJ13105 (FLJ13105), mRNA	1.2
5061		Helicobacter pylori pfs protein homolog (pfs) and sigma 80 (rpoD)	
5261	AF009177	genes, complete cds	0.6
5262	17015150	Arabidopsis thaliana genomic DNA, chromosome 5, TAC	
5262	AB015470	clone:K1B16	5.2
5264	L08970	Nematode choline acetyltransferase (cha-1) gene, exons 1-11 and	
5265		complete cds	0.29
	XM 046294	Brassica rapa DNA for S-receptor kinase, complete cds	0.69
	XM 027078	Homo sapiens testis expressed sequence 15 (TEX15), mRNA	0.21
2207	7LIVI_02/0/8	Homo sapiens ATPase, Class V, type 10D (ATP10D), mRNA	0.22

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		The groot (Blast VS. Genoalik)	
NO	ACCESSN	DESCRIP	P VALUE
		Mus musculus adult male hippocampus cDNA, RIKEN full-length	1 VALUE
5268	AK019309	enriched library, clone:2900009119, full insert sequence	2E-13
5269	NM_016985	Mus musculus myotubularin related protein 1 (Mtmrl), mRNA	0.0004
		Mus musculus 13 days embryo male testis cDNA, RIKEN full-length	0.0004
5270	AK020049	enriched library, clone:6030408K18, full insert sequence	1.8
		and the sequence	1.0
5271	AF323989	Danio rerio transcriptional intermediary factor 2 mRNA, complete cds	6.3
		Pisum sativum trans-cinnamic acid hydroxylase (CYP73A9) gene.	0.5
5272	AF175275	CYP73A9-v1 allele, partial cds	0.073
5274	AF327749	Mus musculus smoothelin gene, promoter and partial sequence	6E-11
5275	AJ276633	Glomus mosseae partial TOR2 gene, exons 1-9	6
		g-ma, vacabit y	
- 1		Boophilus microplus cytochrome b apoenzyme (Cytb) gene, partial	
- 1		cds; tRNA-Ser, tRNA-Leu, tRNA-Cys, and tRNA-Met genes,	
		complete sequence; and NADH dehydrogenase subunit 2 (ND2) gene.	
5276	AF110612	partial cds, mitochondrial genes for mitochondrial products	0.072
5278	XM 043395	Homo sapiens fatty acid desaturase 2 (FADS2), mRNA	3E-36
		Homo sapiens zinc finger protein 133 (clone pHZ-13) (ZNF133),	312-30
5279	XM_046528	mRNA	2
5280	T10000#		
3280	U20807	Bos taurus protein tyrosine phosphatase BA14 mRNA, complete cds	0.19
5281	VM nacaca	Homo sapiens hypothetical gene supported by AK026802	
3261	XM_036368	(LOC91132), mRNA	0.026
5282	XM_007023	Homo sapiens G protein-coupled receptor kinase-interactor 2 (GIT2), mRNA	
3202	AIVI_007023		7E-44
5283	U86362	Saccharomyces cerevisiae MAL gene divergent promoter region, and	
5284	Y11740	hypothetical protein, complete cds	0.24
3204	111/40	H.sapiens whn gene, exon 1a and 1b	0.22
5285	S78915	Office administration of the state of the st	
5200	576913	{Mu1 element insertion site, clone 11} [maize, Transposon, 300 nt]	0,002
5286	AF310896	Dictyostelium discoideum RacJ (racJ) gene, complete cds, LagC-like protein (lagC3) gene, partial cds, and unknown genes	ł
5287	AJ012333	Buchnera aphidicola plasmid pBTc2, trpE and trpG genes	0.23
5288	AF282046	Homo sapiens clone 15qtel_c321bt7 sequence	0.027
5200	7H 202040		4E-30
5289	L25128	Lycopersicon esculentum auxin-induced proteinase inhibitor (ARPI) gene, complete cds	
		Homo sapiens KIAA1467 protein (KIAA1467), mRNA	0.51
	144_019005	Homo sapieus KiAA1407 piotetti (KiAA1407), ilikiva	2E-46
5291	AE002114	Ureaplasma urealyticum section 15 of 59 of the complete genome	0.054
5292	AF208678	Aedes aegypti clone p502, Pony-Aa-B6 MITE repeat region	3.9
5293	AK024656	Homo sapiens cDNA: FLJ21003 fis, clone CAE03685	5E-55
5294	NC_002355	Bombyx mori mitochondrion, complete genome	2.1
5295		Homo sapiens cDNA FLJ13836 fis, clone THYRO1000734	0,008
5296		Chlamydia muridarum, section 70 of 85 of the complete genome	1.9
		, and a second	4

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		(Diasti Vs. Octionik)	
NO	ACCESSN	DESCRIP	P VALU
		DESCRI	PVALU
		Homo sapiens region containing TLS-associated serine-arginine	
		protein 1; TLS-associated serine-arginine protein 2; TLS-associated	
		serine-arginine protein 2; TLS-associated serine-arginine protein 2;	
5297	XM_032347	TLS-associated serine-arginine protein 1; TLS-associa>	0
		H.sapiens CpG island DNA genomic Mse1 fragment, clone 78e4,	
5298	Z63116	reverse read cpg78e4.rt1a	0.24
5299	AE006833	Sulfolobus solfataricus section 192 of 272 of the complete genome	6.4
		Homo sapiens similar to ZINC FINGER PROTEIN 20 (ZINC	0.4
		FINGER PROTEIN KOX13) (DKFZP572P0920) (H. sapiens)	
5300	XM_032811	(LOC90591), mRNA	0.21
		Homo sapiens DNA methyltransferase (DNMT1) gene, exons 2, 3,	0.51
5301	AF202552	and 4	0.23
		Pygathrix roxellana NADH dehydrogenase subunit 3 (ND3) gene,	
		partial cds, tRNA-Arg gene, complete sequence, NADH	
J		dehydrogenase subunit 4L (ND4L) and NADH dehydrogenase subunit	
-		4 (ND4) genes, complete cds, and tRNA-His, tRNA-Ser and tRNA-	
5302	U92963	Leu genes, comp>	0.17
i			
		Euglena deses chloroplast psbC gene: complete group III twintron,	
5303	Z99833	complete internal mat1 gene, partial 5' and 3' psbC exons	4.7
5304	AB047962	Macaca fascicularis brain cDNA, clone:QnpA-13041	e-125
		Euplotes crassus transposon Tecl clone Tecl-2 orf 2 and orf 3	
5305	AF159913	pseudogenes, complete sequence	0.5
5306	Y15435	Kluyveromyces lactis PDC1 gene, promoter region	0.087
5307	U67506	Mothamassana iannasshii sastan 40, 6160, 631	
5507	007300	Methanococcus jannaschii section 48 of 150 of the complete genome Bacillus subtilis phosphofructokinase I (pfkI) gene, partial cds; and	1.4
5308	U73943	pyruvate kinase I gene, complete cds	
500	313373	Bos taurus photoreceptor disk rim specific protein rom-1 (ROM1)	2.2
5309	U72027	mRNA, partial cds	0.06
5310	X16509	Rice alpha-amylase gene	2.1
5312	AB049900	Macaca fascicularis brain cDNA, clone:OnpA-19713	0.01
5313	Z24756	S.pombe rhp51 and rpa1 genes, complete CDS's	0.003
		Homo sapiens similar to hypothetical protein FLJ10546 (H. sapiens)	0.003
5314	XM_002989	(LOC93548), mRNA	5E-62
5316	AL358272	S.pombe chromosome I cosmid c458	0.61
5317	XM_044123	Homo sapiens cadherin 20, type 2 (CDH20), mRNA	4E-48
		Bacteroides fragilis transposon Tn5520 transposase (bipH) and	
5318	AF038866	mobilization protein BmpH (bmpH) genes, complete cds	0.73
	1		
5319	AF332577	Homo sapiens prosomal P27K protein (PSMA6) gene, partial cds	2E-51
5320	X68650	O.cuniculus mRNA for ryanodine receptor	0.55

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AC013455

AJ234722

X79699

AB063075

Z56909

L44121

XM 050246 mRNA

sequence

H.sapiens ALU repeat, 230bp

reverse read cpg152h9,rt1a

0.23

0.24

0.0003

0

2E-19

0.69

0.069

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#### Table 3A Nearest Neighbor (BlastN vs. Genbank) SEQ ID NO ACCESSN DESCRIP P VALUE H.sapiens CpG island DNA genomic Mse1 fragment, clone 69d2, 5322 Z65960 reverse read cpg69d2.rt1b 17 Candida albicans 60S acidic ribosomal protein type P2-A (p2A) gene, 5323 AF317661 complete cds 5.8 5324 Thermotoga maritima section 88 of 136 of the complete genome AE001776 0.64 5325 NM 031987 Rattus norvegicus carnitine octanoyltransferase (COT), mRNA 0.62 5326 AB035500 Rana rugosa gene for FTZ-F1, exon 1 0.009 Methanococcus jannaschii section 32 of 150 of the complete genome 5327 U67490 0.2 Glycine max mRNA for hypothetical protein, complete cds. 5328 AB060000 clone:SSC2 0.22 5329 U50840 Borrelia garinii 70 kbp plasmid D6 protein gene, complete cds 0.072 5330 1167535 Methanococcus jannaschii section 77 of 150 of the complete genome 2.1 AK001125 Homo sapiens cDNA FLJ10263 fis, clone HEMBB1000991 3E-43 Ascogaster sp. 16S ribosomal RNA gene, mitochondrial gene for AF029114 5333 mitochondrial RNA, partial sequence 0.068 5334 AF320594 Homo sapiens PKD1P4 pseudogene, exons 2 through 15 6.3 Oryza sativa microsatellite MRG3819 containing (TA)X27, genomic 5335 AY021494 sequence 0.024 5336 AF146651 Homo sapiens glyoxalase-I gene, complete cds 0.056 5337 AB054516 Globicephala melas DNA, SINE flanking sequence Mago22 locus 0.024 Homo sapiens (subclone 1_f6 from P1 H56) DNA sequence, complete 5338 AC001052 sequence 0.003 5339 U00951 Human clone A9A2BR11 (CAC)n/(GTG)n repeat-containing mRNA 0.65 5340 AF298180 Caenorhabditis elegans tropomyosin isoform IV mRNA, complete cds 0.64 Homo sapiens hypothetical gene supported by AK025398 XM 042336 (LOC92036), mRNA 5341 2.1 L44121 5342 Homo sapiens (clone pHK2.1D) CMT1A gene, repeat unit 0.0003 Escherichia coli O157:H7 EDL933 genome, contig 3 of 3, section 175 5343 AE005556 of 290 0.23 5344 AB035185 Homo sapiens RHD gene, intron 9, complete sequence 0.63 Homo sapiens BAC clone CTD-2347O14 from 7p12-p14, complete

Hordeum vulgare genomic DNA fragment; clone MWG2031.rev

Homo sapiens secretory protein SEC8; KIAA1699 protein (SEC8),

H. sapiens CpG island DNA genomic Mse1 fragment, clone 152h9,

Macaca fascicularis brain cDNA clone:QtrA-11888, full insert

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		garat (a sad the contains)	
NO	ACCESSN	DESCRIP	
		Drosophila melanogaster genomic scaffold 142000013385665,	P VALU
5352	AE002722	complete sequence	
		1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1.5
	l	Stachyarrhena sp. Jansen-Jacobs 4707 ribosomal protein S16 (rps16)	
5353	AF201021	gene, partial intron sequence; chloroplast gene for chloroplast product	
5354	Y17506	Euplotes octocarinatus fRNA-Cys gene	
		T stockamand indus-cys gene	0.024
5356	XM 040513	Homo sapiens hypothetical protein FLJ14454 (FLJ14454), mRNA	
		Mus musculus DREAM/calsenilin/KChIP3 gene, complete cds and	0.7
5357	AF287736	exons 3, 4 and 5, alternatively spliced	_
		Danio rerio PITP-less RdgB-like protein (plrdgB) mRNA, complete	2
5358	AF230807	cds	
			6,1
5359	U67563	Methanococcus jannaschii section 105 of 150 of the complete genome	
		Methanobacterium thermoautotrophicum from bases 1655364 to	0.24
5360	AE000935	1666496 (section 141 of 148) of the complete genome	
		Rattus norvegicus FAT tumor suppressor (Drosophila) homolog (Fat),	0.21
5361	NM 031819	mRNA	
5362	XM 005889	Homo sapiens kinesin-like 1 (KNSL1), mRNA	5,2
5363	AF339829	Homo sapiens clone IMAGE:609847, mRNA sequence	3.6
-	12007027	Mus musculus circadian locomoter output cycles kaput (Clock),	0.022
5364	NM 007715	mRNA (Clock),	
		Homo sapiens hypothetical protein DKFZp586F1122 similar to	e-116
5365	XM 016117	axotrophin (DKFZP586F1122), mRNA	
_		Homo sapiens regulator of differentiation (in S. pombe) 1 (ROD1),	0
5366	XM_005554	mRNA (ROD1),	
_		III.COM	6.3
5368	X76311	H. sapiens endothelial nitric oxidase synthase gene, exons 15 and 16	
		Homo sapiens proteoglycan 4, (megakaryocyte stimulating factor,	0.7
5369	XM_001738	articular superficial zone protein) (PRG4), mRNA	
<del></del>		Methanobacterium thermoautotrophicum from bases 1655364 to	0.63
5370	AE000935	1666496 (section 141 of 148) of the complete genome	
5371	Y18011	Phaedon cochleariae mRNA for chitinase	0.23
5372	Z99768	Flaveria trinervia gdcsP pseudogene	0.021
		Fusobacterium ulcerans strain NCTC 12112 16S ribosomal RNA	4.5
- 1		gene, partial sequence; internal transcribed spacer 1, complete	
5373	AF342852	sequence; and 23S ribosomal RNA gene, partial sequence	
		Homo saniene hypothetical perteir El moses	0.18
5374	BC003397	Homo sapiens, hypothetical protein FLJ20505, clone MGC:4960 IMAGE:3448518, mRNA, complete cds	
		Homo caniona MODA OTODA	1E-69
5375	AF385088	Homo sapiens NOD2 (NOD2) gene, NOD2-2722G>C allele, exon 8 and partial cds	
		Papio cynocephalus provirus, complete genome	0.58
		Rattus populatus provintus, complete genome	0.22
377	AF121217	Rattus norvegicus pro-alpha-2(I) collagen (col1a2) mRNA, complete	
-			5.5
378	BC009075	Mus musculus, beta-1,3-N-acetylglucosaminyltransferase 1, clone	- 1
		MGC:6892 IMAGE:2654354, mRNA, complete cds	1.8

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	1	Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ II		Total Cot Picignoon (Diastry Vs. Gendank)	
NO	ACCESSN	DESCRIP	1
	1	DESCRIP	P VALUE
5379	XM 00134	1 Homo sapiens DKFZP547E1010 protein (DKFZP547E1010), mRNA	
5380	XM_04640	8 Homo sapiens KIAA1041 protein (KIAA1041), mRNA	3E-15
		Mus musculus 6 days neonate skin cDNA, RIKEN full-length	0.52
5381	AK020701	enriched library, clone: A030009B12, full insert sequence	
		actual, cione.7030009B12, full liseri sequence	0.000001
5383	U32711	Haemophilus influenzae Rd section 26 of 163 of the complete genome	
		20 of 103 of the complete genome	0.021
5384	XM 046598	Homo sapiens DKFZP434K156 protein (DKFZP434K156), mRNA	
		15-44150 protein (DKF2F454K150), miknA	0.58
	]	Buchnera aphidicola anthranilate synthase large subunit (trpE) gene	
5385	L43551	and anthranilate synthase small subunit (trpG) gene, complete cds	
5386	XM 012233	Homo sapiens MDS023 protein (MDS023), mRNA	0.65
5387	AF139358	Homo sapiens BUB1 protein gene, exon 17	0.6
		Mus musculus adult retina cDNA, RIKEN full-length enriched	0.008
5388	AK020916	library, clone:A93003 IL14, full insert sequence	
		Homo sapiens mRNA; cDNA DKFZp547E184 (from clone	1.5
5389	AL390170	DKFZp547E184)	1.0
5390	AK021659	Homo sapiens cDNA FLJ11597 fis, clone HEMBA1003856	0.21
5391	AK024200	Homo sapiens cDNA FLJ14138 fis, clone MAMMA1002765	0.00003
5392	AJ011727	Homo sapiens partial TOP2 beta gene, sequence G	0.00003
		Homo sapiens leukemia inhibitory factor receptor (LIFG) gene,	0.00003
5393	AF018079	alternative promoter which is functional in non-placental tissues	6.2
		Macaca fascicularis brain cDNA clone:QtrA-14876, full insert	0.2
5394	AB063085	sequence	0.026
		Homo sapiens olfactory receptor HPFH1OR (HPFH1OR) gene,	0.020
5395	AF154673	complete cds	0.23
			0.23
5396	AE007505	Streptococcus pneumoniae section 188 of 194 of the complete genome	6.6
5397	Z75199	S.cerevisiae chromosome XV reading frame ORF YOR291w	0.081
- 1		Bombyx mori Pao-like retrotransposon Kamikaze DNA similar to	0.001
5398	AB042120	polyprotein coding region	0.24
5399	AF379854	Cloning vector pVLH/hsp, complete sequence	0.0003
5400	XM_009648	Homo sapiens HSPC072 protein (HSPC072), mRNA	0.022
5401	AB056333	Macaca fascicularis brain cDNA, clone:OflA-13486	9E-14
5402	AK021824	Homo sapiens cDNA FLJ11762 fis, clone HEMBA1005670	3E-46
		Human DNA sequence from clone RP11-30M17 on chromosome 6.	
5403	AL513010	complete sequence [Homo sapiens]	0.026
5404	AK023753	Homo sapiens cDNA FLJ13691 fis, clone PLACE2000100	0.077
ĺ		Mus musculus 18 days pregnant adult female placenta and extra	
		embryonic tissue cDNA, RIKEN full-length enriched library	
5406	AK014452	clone:3830422K02, full insert sequence	0.17
5407	XM_003220	Homo sapiens polybromo 1 (PB1), mRNA	1E-35
5408	770010		
7406	Z78918	H.sapiens flow-sorted chromosome 6 HindIII fragment, SC6pA19C6	1.5

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	1	Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ II	,	Table 52 Nearest Neighbor (Blastin vs. Genbank)	_
NO	ACCESSN		ł
NO	ACCESSIV		P V
	1	Mycoplasma pirum purine phosphorylase gene, 3' end,	
	1	deoxyriboaldolase, thymidine phosphorylase, cytidine dcaminase,	
5409	V 12200	phosphomannomutase genes, complete cds, and triose phosphate	
	L13289	isomerase gene, 5' end	0
5410	AC024203	Caenorhabditis elegans cosmid Y72A2A, complete sequence	(
		Homo sapiens translocation breakpoint MLLT1/MLL fusion gene.	
5411	AF373586	partial sequence	0
		A pleuropneumoniae tfbA gene for transferrin-binding protein (1949	_
5412	Z46774	[bp)·	0.
		Methanosarcina acetivorans strain C2A MtaF (mtaF) and MtaG	-
5413	AF319044	(mtaG) genes, complete cds	0.
		Deinococcus radiodurans R1 section 70 of 229 of the complete	
5414	AE001933	chromosome 1	6
		Human platelet-derived growth factor A type receptor mRNA,	
5415	M22734	complete cds	_
		Asterella tenella trnL gene, intron sequence; chloroplast gene for	0.
5416	AF264650	chloroplast product	١
		Sambucus nigra clone VBL85 thaumatin-like protein (TLPI) mRNA,	6
5417	AF378571	complete cds	
0.12.	111 5 7 65 7 1	comprete cas	1
5418	U53183	Pottus namunian 250 LD	
5419	AK026673	Rattus norvegicus 250 kDa estrous-specific protein mRNA, partial cds	0.
3419	AK0200/3	Homo sapiens cDNA: FLJ23020 fis, clone LNG00943	0.2
5420	<b>TD ( 04104</b>	Homo sapiens NADH dehydrogenase (ubiquinone) 1 beta subcomplex,	
3420	XM_041367	4 (15kD, B15) (NDUFB4), mRNA	2E
		Arabidopsis thaliana unknown protein (F11F12.23) mRNA, complete	
5421	AY039939	cds	5.
[		Mus musculus splicing factor Sc35 (Pr264) mRNA, partial eds,	
5422	AF250135	alternatively spliced	2.
5423	M32328	C.pipiens esterase B1 gene, complete cds	0.0
- 1		Bos taurus inward rectifier potassium channel BIK mRNA, complete	
5424	U95369	cds	0.0
T			
5425	AE002157	Ureaplasma urealyticum section 58 of 59 of the complete genome	5.1
5426	AF270633	Homo sapiens clone 5qtel c25t3 sequence	9E-
5427	Z28252	S.cerevisiae chromosome XI reading frame ORF YKR027w	0.1
		Drosophila melanogaster genomic scaffold 142000013385253,	0.1
5428	AE002601	complete sequence	
5429	AJ403990	Hyla arborea microsatellite DNA, clone WHA1-61	0.2
		Rattus norvegicus protein tyrosine phosphatase, non-receptor type 5	3.8
5430	NM_019253	(Ptpn5), mRNA	
			0.7
5431	AE006543	Streptococcus pyogenes M1 GAS strain SF370, section 72 of 167 of	
		the complete genome	0.6
5432	AFILICACE	Homo sapiens islet cell autoantigen ICA69 (ICA1) gene, exon 1 and	
5433	AF146363	partial cds	0.07
	M96943	Human profilaggrin gene exons 1-3, 5' end	0.06
5434	AK025246	Homo sapiens cDNA: FLJ21593 fis, clone COL07050	0.13

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		The second secon	т—
NO	ACCESSN	DESCRIP	P VALUE
		Homo sapiens general transcription factor IIIC, polypeptide 4 (90kD)	1 11202
5435	XM_027835	(GTF3C4), mRNA	5.9
5437	AJ223441	Branchiostoma floridae Pax6 gene (AmphiPax6), clone J2	0.24
5438	AB048345	Equus caballus DNA, chromosome Xp21, microsatellite TKY39	0.73
5439	AB049844	Macaca fascicularis brain cDNA, clone:QnpA-18434	2.2
5440	TD 7 00 50 10	Homo sapiens similar to nuclear pore membrane glycoprotein 210 (M.	
5440	XM_036740		0.19
5441	AFC010527	Mus musculus adult male testis cDNA, RIKEN full-length enriched	
3441	AK019537	library, clone:4921505C17, full insert sequence	0.081
5442	AE003964	Xylella fastidiosa 9a5c, section 110 of 229 of the complete genome	6
5443	X95276	P.falciparum complete gene map of plastid-like DNA (IR-B)	0.008
5444	AF310264	Ovis aries leptin mRNA, partial 3' UTR	2.2
5445	XM_048688		2.2
- 1		Arabidopsis thaliana chromosome II section 209 of 255 of the	
5446	AC007133	complete sequence. Sequence from clones T6A23, F13I13	2
5447	AJ011727	Homo sapiens partial TOP2 beta gene, sequence G	0.00003
		Homo sapiens cAMP responsive element modulator (CREM) gene,	
5448	AF214665	exon H	0.52
****	4 770 5004 -	Staphylococcus epidermidis strain SR1 clone step.1002a02 genomic	
5449	AF269319	sequence	1.9
5450	4 E001 202	Plasmodium falciparum chromosome 2, section 30 of 73 of the	
3430	AE001393	complete sequence	0.72
5451	XM 002437	Homo sapiens cAMP-regulated guanine nucleotide exchange factor II (CAMP-GEFII), mRNA	
5452	AF298224	Homo sapiens RPCI-II 289E13 CD21 gene, partial cds	7E-32
5453	AF254641	Homo sapiens oxytocin/vasopressin intergenic region	0.007
0.00	111 25 10 11	Homo sapiens phosphate cytidylyltransferase 1, choline, beta isoform	0.01
5454	XM_010303	(PCYT1B), mRNA	2
		Mus musculus 18 days pregnant adult female placenta and extra	
- 1		embryonic tissue cDNA, RIKEN full-length enriched library,	
5455	AK014447	clone:3830421F13, full insert sequence	0.22
		P.hybrida 5-enolpyruvylshikimate 3-phosphate synthase mRNA,	
5456	M21084	complete cds	0,24
		Homo sapiens Microfibril-associated glycoprotein-2 (MAGP2),	
5457	XM_006648	mRNA	1.4
		Gallus gallus T cell receptor delta chain (TCRD) pseudogene, partial	
5458	AF175434	sequence	0.76
5459	AE000775	Aquifex acolicus section 107 of 109 of the complete genome	0.0008
5460	VA. 0212C	Homo sapiens similar to KIAA0377 gene product (H. sapiens)	
2400	XM_031261	(LOC90371), mRNA	1.6
5461	BC008389	Homo sapiens, Similar to hypothetical protein LOC57821, clone IMAGE:4076369, mRNA	
2401	DC000309	191AGE.4070309, MKNA	0.24

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		Jan ( and  and  and  and  and  and  and	т —
NO	ACCESSN	DESCRIP	P VALUE
i		Streptococcus thermophilus Peb1 (peb1), GlnQ (glnQ), response	
		regulator Rr1 (rr1), Hpk2 (hpk2), YycJ-like protein (vycJ), and MurM	ĺ
5462	AF327739	(murivi) genes, complete cds	0.63
		HIV-1 isolate NC3940-1998 from USA pol polyprotein (pol) gene,	
5463	AY030939	partial cds	0.026
5464	XM_049237		0.053
5465	4.770.471.4	Paracheilognathus imberbis mitochondrial control region, partial	
5466	AY017147 X81451	sequence	0.25
5467	AF271036	N.frontalis enolase gene, promotor region	0.009
3407	AF2/1030	Mus musculus clone rF12m odorant receptor gene, partial cds	2E-08
5468	J05637	Yeast (S.cerevisiae) 70 kDa heat shock protein (SSA4) gene, complete	
5469	X75420	P.falciparum (FCR3) cpn60 gcne	0.22
	2073420	r Jaiciparum (FCR3) cpnou gene	0.025
		A sinotoho stan aslamati - VIIIDao - il I van an	
		Acinetobacter calcoaceticus KHP18 partial pKLH2 plasmid including	
5470	AF213017	aberrant mercury resistance transposon TnPKLH2, truncated insertion sequence IS 1011.D1, and determinants for CinH resolution system	
5471	AC024806	Caenorhabditis elegans cosmid Y52E8A, complete sequence	1.9
	110021000	Drosophila melanogaster filamin isoforms A and B (Filamin) gene,	0.026
5472	AF183178	lexons 1 and 2	0.043
			0.043
5473	AF209925	Plasmodium falciparum STARP antigen (STARP) gene, complete cds	2.1
		Ectomycorrhizal isolate ECM 133 internal partial internal transcribed	
- 1		spacer 1 (ITS1), 5.8S ribosomal RNA gene, and partial internal	
5474	AJ410864	transcribed spacer 2 (ITS2); isolate ECM 133	2
5475	NC_001889	Dictyostelium discoideum plasmid Ddp5, complete sequence	2.1
		Boophilus microplus clone PNC 189 isolate Kilkivan2 microsatellite	
5476	AY009594	sequence	0.19
5477	U55723	Human ataxia-telangiectasia (ATM) gene, exon 25	0.69
5.50		Saguinus oedipus MHC class II antigen (Saoe-DRB1) gene, Saoe-	
5478 5479	AF197229 AE001290	DRB1*0303 allele, partial sequence	0.00004
34/9	AE001290	Chlamydia trachomatis section 17 of 87 of the complete genome	0.2
5480	XM 016611	Homo sapiens similar to 8-oxoguanine DNA glycosylase (H. sapiens) (LOC93577), mRNA	
3400	AW_010011	CLOC933//), mRNA	0.069
5481	AJ224148	Saccharomyces cerevisiae mitochondrial Tyr-tRNA, Asn-tRNA and Mct-tRNA genes	
5482	AK022243	Homo sapiens cDNA FLJ12181 fis, clone MAMMA1000746	0.026
		Legionella pneumophila Philadelphia-1 intracellular multiplication	0.17
5483	U07354	region (icmV, icmW, icmX, lphB) genes, complete cds	
		Plasmodium falciparum chromosome 2, section 3 of 73 of the	0.33
5484	AE001366	complete sequence	0.14
		Plasmodium falciparum chromosome 2, section 22 of 73 of the	0.14
5485	AE001385	complete sequence	1.5
		Homo sapiens (subclone 1_a3 from P1 H54) DNA sequence, complete	1.3
5486	L81686	sequence	0.62
5487	XM_017648	Homo sapiens leucine-rich repeat-containing 2 (LRRC2), mRNA	0.0006
			/

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-	_		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEC	ŠΒ			
14	<u>-  </u>	ACCES		DYZAY
54	。 l	AE0013	Plasmodium falciparum chromosome 2, section 9 of 73 of the	P VAL
548		Z46943		1.9
549		U68246		0.58
34,	~+	008240	Dictyostelium discoidenm Wac A (woo A)	
549	01	AK00995		0.00.
	+	ALKOUSS.		0.08
549	12	AF19079	Arabidopsis thaliana subtilisin-type serine endopeptidase XSPI	0.00
	-	111 15075		1.9
549	3	AF33679	Plasmodium falciparum dynamin-like protein (dyn) mRNA, complete	1
	+	14 55077	cus	9.6
549	4	AB03584	4 Coturniy coturniy in	
	+			0.026
5495	5	AF127240		
	_			0.42
5496	5   .	AY019789	Oryza sativa microsatellite MRG2114 containing (AT)X78, genomic sequence	
5497		AF106953		0.003
5498		M_00717		0.43
	$\top$			0.014
5499	N	M 00691	Homo sapiens retinitis pigmentosa 2 (X-linked recessive) (RP2),	
5500	7	X95961	A.thaliana CER3-like gene	0.003
	T		Homo sapiens inositol 1,4,5-triphosphate receptor, type 1 (ITPR1),	0.071
5501		M_041428		
5502		S50200	dopamine beta-hydroxylase [mice, Genomic/mRNA, 2274 nt]	0.18
5503		E001178		2.1
5504		Z74797		0.026
5505		X15979	Plasmodium falciparum alpha-tubulin I gene	2.1
	1			0.0008
506	I A	F221538	Cryptosporidium felis heat shock protein 70 (HSP70) gene, partial cds	
	1.			1.8
507	I A	F083226	complete cds	0.00
508	١			0.69
508	A	F262729	Ornithorhynchus anatinus olfactory receptor pla32 gene, partial cds	6E-10
	ı			0E-10
509	١.,	263057	1 Sounds, complete cus; milochondrial gange for mit-1 - 1 - 1	
	A	203037		2
- 1			Agrocybe aegerita B type DNA polymerase (Mtpol) gene, complete	
510	ΔΤ	061244		
11		64612		0.003
-			H.sapiens dystrophin-gene (introns 13/42)	0.69
12	AF	269472	Staphylococcus epidermidis strain SR1 clone step.1005a12 genomic sequence	
14			Homo sapiens GE36 gene, exon 8	0.21
15		391744	S.pombe chromosome I cosmid c17D4	0.12
7			Homo seriese risella d	0.42
16	XМ	034682	Homo sapiens similar to general transcription factor II, i (H. sapiens) (LOC90875), mRNA	
	_			7E-22

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	-	Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
5517	AF361817	Arabidopsis thaliana At1g06730/F4H5_22 gene, complete cds	0.69
5518	Z48243	A.thaliana PARP mRNA for PARP protein	0.69
5519	AL390169	Homo sapiens mRNA; cDNA DKFZp547D064 (from clone DKFZp547D064)	- 100
	122010107	Homo sapiens hypothetical protein DKFZp434K1421	e-180
5520	NM_032141	(DKFZP434K1421), mRNA	e-169
5521	XM 037214	Homo sapiens laminin, alpha 5 (LAMA5), mRNA	0.62
		Botrytis cinerea strain T4 cDNA library under conditions of nitrogen	
5522	AL110675	deprivation	0.008
		Dasyurus viverrimus/maculatus mixed DNA library microsatellite	
5523	AF124216	Q4.4.10 sequence	1.9
		Human chromosome 14 DNA sequence Partial sequence from BAC R-	
		1109N18_PCR1 of library RPCI-11 from chromosome 14 of Homo	
5524	AL583762	sapiens (Human), complete sequence	6E-42
5525	AC024758	Caenorhabditis elegans cosmid Y37E11AM, complete sequence	1.4
		Homo sapiens MER receptor tyrosine kinase (MERTK) gene, exons	
5526	AF260528	17 and 18	3E-35
		Plasmodium falciparum malaria exported protein-1 (EXP-1) gene.	
5527	AF246997	complete cds	0.019
		Homo sapiens sigma receptor (SR31747 binding protein 1) (SR-BP1),	
5528	XM 011810	mRNA	3.9
5529	XM_011100	Homo sapiens complement component 7 (C7), mRNA	5.9
5530	X64315	S.scrofa mRNA for Na/D-glucose cotransporter regulatory-subunit	0.0007
		Homo sapiens bromodomain adjacent to zinc finger domain, 2A	
5531	XM_048945	(BAZ2A), mRNA	3.4
5532		Homo sapiens DKFZP566I1024 protein (DKFZP566I1024), mRNA	1E-10
5533	XM_002977	Homo sapiens ring finger protein 13 (RNF13), mRNA	2.8
	1 72 00 2 5 0		
5534	AJ300359	Homo sapiens partial RYR2 gene for ryanodine receptor 2, exon 20	1E-36
5535	BC008590	Homo sapiens, hypothetical protein FLJ21313, clone MGC:16820 IMAGE:4148772, mRNA, complete cds	e-154
3333	BC008370	Homo sapiens secretory protein SEC8; KIAA1699 protein (SEC8),	e-134
5536	XM_050246	mRNA (SECS),	0
		Homo sapiens mRNA; cDNA DKFZp547E024 (from clone	
5537	AL442094	DKFZp547E024)	2
1 1		Tribolium castaneum Tc-tailless gene, upstream regulatory region and	
5538	AF225975	partial cds; tRNA-Tyr gene, complete sequence; and unknown gene	0.24
1 1		Sidalcea malviflora x Sidalcea oregana 5.8S rRNA gene, internal	
5539	AJ304934	transcribed spacer 1 (ITS1) and internal transcribed spacer 2 (ITS2)	2.1
		Helicobacter pylori, strain J99 section 89 of 132 of the complete	
5540	AE001528	genome	0.009

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Homo sapiens putative DNA/chromatin binding motif (PLU-1),	
5541	XM_046698	mRNA	2.5
5542	AF101309	Caenorhabditis elegans cosmid H24G06, complete sequence	2.1
5543	AL023839	Caenorhabditis elegans cosmid Y39A1C, complete sequence	1.4
5544	XM_017673	Homo sapiens hypothetical protein FLJ21934 (FLJ21934), mRNA	0
		Homo sapiens 11 beta-hydroxysteroid dehydrogenase 2 (HSD11B2)	
5545	U27317	gene, complete cds	2E-92
5546	XM_045095	Homo sapiens KIAA1320 protein (KIAA1320), mRNA	0.6
		Lepidopilum scabrisetum ribosomal protein system 4 (rps4) gene,	
5547	AF143066	partial cds	0.25
		Homo sapiens, Similar to dihydropyrimidinase-like 2, clone	
5548	BC010880	MGC:1757 IMAGE:3542012, mRNA, complete cds	5.2
-		Lycopersicon esculentum auxin-induced proteinase inhibitor (ARPI)	
5549	L25128	gene, complete cds	0.55
5550	NM_030665	Homo sapiens retinoic acid induced 1 (RAI1), mRNA	9E-57
5552	AF127699	Erythroxylum argentinum chloroplast atpB-rbcL spacer	5.5
5553	U97192	Caenorhabditis elegans cosmid C01F4, complete sequence	2.7
		Arabidopsis thaliana 14-3-3 protein GF14 epsilon (GRF10) gene,	
5554	AF145302	complete cds	8.9
		Citrus sinensis capsanthin/capsorubin synthase (CCS) gene, complete	
5555	AF169241	cds	0.21
		Tetrahymena thermophila dynein heavy chain (DYH6) gene, partial	
5556	AF153268	cds	1E-15
5557	L13966	Mouse delta/YY1/NF-E1/UCRBP transcription factor, exon 3	0.65
		Streptococcus gordonii Sagp-like protein gene, complete cds; and Oct-	
5558	AF172730	like protein gene, partial cds	1.9
		Human DNA sequence from clone RP11-124P3 on chromosome X,	
5559	AL591422	complete sequence [Homo sapiens]	1.8
5560	AF328433	Homo sapiens clone 2qtel_c275_24t3 sequence	4.8
		Podarcis pityusensis cytochrome b (cytb) gene, mitochondrial gene	
5561	AF052640	encoding mitochondrial protein, partial cds	2.3
		Cicindela marutha 16S large subunit ribosomal RNA, mitochondrial	
5562	AF133000	gene for mitochondrial product, complete sequence	0.024
5563	AJ239057	Antirrhinum majus far gene for farinelli protein	5,1
5564	AL032651	Caenorhabditis elegans cosmid Y6D1A, complete sequence	2
		Homo sapiens partial SNAP-23 gene for synaptosome associated	
5565	AJ278974	protein-23, exons 6-8	1.8
5566	M33379	Guinea pig lipoprotein lipase (gplpl) gene, exon 3-7	0.46
		Rheum reticulatum tRNA-Leu (trnL) gene and trnL-trnF intergenic	
5567	AF303443	spacer, partial sequence; chloroplast gene for chloroplast product	0.018
		Homo sapiens (subclone 1_g10 from P1 H39) DNA sequence,	
5568	L81826	complete sequence	4E-16
5569	NM_005958	Homo sapiens melatonin receptor 1A (MTNR1A), mRNA	2
5570	AJ133356	Homo sapiens MSMB gene (partial), exons 2-3	0.66

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		Table 511 Holicat Heighbor (Diases Va. Octobrik)	
NO	ACCESSN	DESCRIP	P VALUE
		Human herpesvirus type 7 U38 gene, U39 gene for glycoprotein B and	
5571	D86577	U40 gene for DNA polymerase, complete and partial cds	0.5
		Streptococcus pyogenes M1 GAS strain SF370, section 107 of 167 of	
5572	AE006578	the complete genome	0.008
5573	U31170	Streptococcus mutans ATPase operon, complete sequence	0.041
		Schistosoma mansoni thioredoxin peroxidase 3 (TPx3) gene, complete	
5575	AF301004	cds	0.23
		Staphylococcus epidermidis strain SR1 clone step. 1000d09 genomic	
5576	AF269295	sequence	2
	1.F11.400.4	Saccharomyces sp. S6U ATP synthase subunit 8 (ATP8) gene,	
5577 5578	AF114934 AF261143	mitochondrial gene encoding mitochondrial protein, complete cds  Homo sapiens chromosome 9 lung cancer-associated mRNA	0.008
5579	X95276	P.falciparum complete gene map of plastid-like DNA (IR-B)	2
5580	NM 030844	Rattus norvegicus islet cell autoantigen 1, 69 kDa (Ica1), mRNA	0,00003
3380	NW_030844	Homo sapiens proteoglycan 4, (megakaryocyte stimulating factor,	0.63
5581	XM 001738	articular superficial zone protein) (PRG4), mRNA	0.21
3361	AIVI_001/38	Macaca fascicularis brain cDNA clone:OtrA-11624, full insert	0.21
5582	AB060860	sequence	5.1
5583	NC 001566	Apis mellifera ligustica mitochondrion, complete genome	0.15
3303	140_001300	Polycycnis ornata tRNA-Leu (trnL) gene, partial sequence; and trnL-F	0.13
		intergenic spacer region, complete sequence; chloroplast gene for	
5584	AF239566	chloroplast product	0.008
5585	X92893	C.roseus GGPP synthase gene	1.6
5586	XM 002601	Homo sapicas desmin (DES), mRNA	5.7
		Rattus norvegicus CD94 antigen (located within the rat natural killer	
5587	NM_012745	gene complex) (Kird1), mRNA	1.4
5588	AK022035	Homo sapiens cDNA FLJ11973 fis, clone HEMBB1001221	0.023
		Plasmodium falciparum 1-CysPxn mRNA for 1-cys peroxidoxin,	
5589	AB020595	complete cds	1.4
	i	Human DNA sequence from clone RP5-839B11 on chromosome 20.	
	ŀ	Contains the first coding exon of the gene for a novel protein with a	
		Kunitz/Bovine pancreatic trypsin inhibitor domain and WAP-type	
5590	AL121778	(Whey Acidic Protein) 'four-disulfide core' domains and an €>	0.07
		Rattus norvegicus voltage gated N-type calcium channel alpha1B	
5591	AF222338	gene, partial cds, alternatively spliced	5.8
5592	AF247193	Mus musculus endobrevin (Vamp8) gene, exon 1	0.18
5593	AC026061	Homo sapiens BAC clone RP11-223K9 from Y, complete sequence	e-163
		Homo sapiens ubiquitin specific protease 9, X chromosome	
5594	XM 030132	(Drosophila fat facets related) (USP9X), mRNA	0.21
<u> </u>		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
5595			
0090	AK015052	library, clone;4930402K13, full insert sequence	0.22
5596	AK015052 AK023333	library, clone;4930402K13, full insert sequence Homo sapiens cDNA FLJ13271 fis, clone OVARC1001000	0.22

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
5599	XM_044123	Homo sapiens cadherin 20, type 2 (CDH20), mRNA	4E-48
5600	XM_030272	Homo sapiens LOC88835 (LOC88835), mRNA	0.3
		Brugia malayi microfilarial sheath protein SHP3a (Bmshp3a) and	
	.=	microfilarial sheath protein SHP3 precursor (Bmshp3) genes,	
5601	AF030944	complete cds	0.089
5602	U59882	Mus musculus MLH1 gene, exon 2	5.2
FC02	47 501 400	Human DNA sequence from clone RP11-113L12 on chromosome 13,	
5603 5604	AL591498 AB002169	complete sequence [Homo sapiens]  Rattus norvegicus RT1.P1 pseudogene for TL antigen	1.8
3004	AB002109	Rattus norvegicus KTT.PT pseudogene for TL antigen	
5605	AB009907	Luciola kuroiwae mitochondrial DNA for 16S rRNA, partial sequence	0.003
5606	AF336829	Rattus norvegicus NEDD8-activating enzyme mRNA, complete cds	0.58
3000	711 330023	Scyliorhimus canicula partial mRNA for follicle stimulating hormone	0.58
5607	AJ310344	beta subunit (fsh beta gene)	0.22
5007	12010011	Does Discourse (said Does Bosso)	- 0.22
5608	D00653	Homo sapiens gene for enteric smooth muscle gamma-actin, exon 8	2
5609	AF203972	Shuttle vector pBA complete sequence	2
		Mus musculus adult male lung cDNA, RIKEN full-length enriched	
5610	AK004804	library, clone:1200016B11, full insert sequence	5.3
		Staphylococcus epidermidis strain SR1 clone step.1040e03 genomic	
5611	AF270009	sequence	0.037
5612	BC006448	Homo sapiens, clone IMAGE:3637085, mRNA	1.9
		Dictyostelium discoideum putative transposon DDT-B, complete	· ·
5613	AF298202	sequence	0.019
5614	AF272001	Ebola virus subtype Zaire strain Mayinga complete genome	0.002
5615	M86544	Cow prostaglandin F synthetase II (PGFSII) mRNA, complete cds	0.075
5616	XM_039684	Homo sapiens putative G-protein coupled receptor (SH120), mRNA	0.16
	4 7200200	Medicago truncatula mRNA for putative AUX1-like permease (lax3	
5617	AJ299399	gene)	6.4
5618	L39214	Ovis aries glucose transporter type 3 (GLUT-3) mRNA, complete cds	0.23
5619	XM 010126	Homo sapiens stromal antigen 2 (STAG2), mRNA	e-127
		H. sapiens (D17S931) DNA segment containing (CA) repeat; clone	
5620	Z23801	AFM248tg5; single read	5E-19
5621	AF210842	Homo sapiens HARP (HARP) gene, exon 17 and complete cds	0.65
		Staphylococcus aureus DNA, complete structure of cassette	
5622	AB047239	chromosome(SCC)-like element, strain:ATCC25923	0.25
5623	AB035193	Homo sapiens RHCE gene, intron 7, complete sequence	2.1
5624	XM_045035	Homo sapiens hypothetical protein FLJ21343 (FLJ21343), mRNA	2.4
5625	XM 052223	Homo sapiens DKFZP564G092 protein (DKFZP564G092), mRNA	0.67
5626	Z28007	S.cerevisiae chromosome XI reading frame ORF YKL007w	0.23

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
5627	M14080	Herpesvirus saimiri thymidylate synthase gene, complete cds	0.025
5628	AF067218	Cacnorhabditis elegans cosmid K10E9	0.071
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	i
5629	AK017102	library, clone:4933437K13, full insert sequence	0.76
5630	XM_038524	Homo sapiens hypothetical protein FLJ23059 (FLJ23059), mRNA	e-118
5631	AB052773	Bombyx mori Bmdsx gene, exon 1, 2, 3, 4, female-specific splicing product complete cds	0.24
5632	AF233069	Galdieria sulphururia maturase (matk) gene, partial eds; 50S ribosomal protein, ribulose-1,3-bisphosphate carboxylase/oxygenase large subunit (rbc.), ribulose-1,5-bisphosphate carboxylase/oxygenase small subunit (rbc.), acetolydroxy-acid synthase small>	0,55
5633	AK027218	Homo sapicns cDNA: FLJ23565 fis, clone LNG10846	0.33
	111027210	HIV-1 isolate WM710-5 from Australia envelope glycoprotein (env)	0.24
5634	AF198462	gene, partial cds	0.61
5636	AF213717	Euhadra quaesita haplotype Mana-B 16S ribosomal RNA gene, partial sequence; mitochondrial gene for mitochondrial product	0.21
5637	AE002730	Drosophila melanogaster genomic scaffold 142000013385730, complete sequence	1.4
5638	AF110027	Candida albicans ATP-dependent transporter (YCF) gene, complete cds	0.069
5639	Z73978	Caenorhabditis elegans cosmid ZC302, complete sequence	0.21
5640	AF379877	Aclista sp. M226 16S ribosomal RNA gene, partial sequence; mitochondrial gene for mitochondrial product	
5641	AF364544	Larix lyallii microsatellite LLY13 sequence	0.52
3041	20.304344	Dermatophagoides farinae allergen Def f II precursor, mRNA, partial	2.2
5642	AF346905	cds	0.008
		Plasmodium falciparum chromosome 2, section 63 of 73 of the	
5643	AE001426	complete sequence	0.001
5644	XM_002850	Homo sapiens TU3A protein (TU3A), mRNA	0.078
5645	AB032899	Rattus norvegicus PIPK2 alpha mRNA for phosphatidylinositol 5- phosphate 4-kinase alpha, complete cds	
3043	AB032699		4.4
5646	AK017370	Mus musculus 6 days neonate head cDNA, RIKEN full-length enriched library, clone:5430431E21, full insert sequence	6E-64
		Homo sapiens similar to GTP-RHO BINDING PROTEIN 1	OD U
5647	XM_042126	(RHOPHILIN) (M. musculus) (LOC91997), mRNA	3E-47
5648	XM_048666	Homo sapiens runt-related transcription factor 1 (acute myeloid leukemia 1; aml1 oncogene) (RUNX1), mRNA	6.3
5649	AJ000259	Caenorhabditis elegans osm-6 mRNA	0.6
5650	AK017713	Mus musculus 8 days embryo cDNA, RIKEN full-length enriched library, clone:5730488L07, full insert sequence	0.087
5651	XM_010294	Homo sapiens arylsulfatase E (chondrodysplasia punctata 1) (ARSE), mRNA	8E-36
5652	XM 039942	Homo sapiens Meis (mouse) homolog 3 (MEIS3), mRNA	0.53

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	0	Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
ИО	ACCESSN	DESCRIP	P VALUE
		Mus musculus 0 day neonate head cDNA, RIKEN full-length enriched	
5653	AK019524	library, clone:4833439O17, full insert sequence	1.7
5654	XM_045122	Homo sapiens KIAA1451 protein (KIAA1451), mRNA	5E-38
		Mus musculus adult male tongue cDNA, RIKEN full-length enriched	
5656	AK009345	library, clone:2310014M14, full insert sequence	3E-35
5657	AF192496	Homo sapicns PDX-1 gene, 5' flanking region	1.5
5658	XM_052620	Homo sapiens KIAA0935 protein (KIAA0935), mRNA	0.0003
5659	XM_049514	Homo sapiens hypothetical protein FLJ13153 (FLJ13153), mRNA	5.6
5660	AJ400965	Proteus mirabilis ORF1 DNA, katA gene and menE gene, strain PR	6.5
		Rattus norvegicus cyclic nucleotide-gated channel beta subunit 1	
5661	NM_031809	(Cngb1), mRNA	0.63
l i		Notothenia coriiceps alphaNCP1 (alphaNCPI), betaNCP1 (betaNCPI),	
		alphaNCP2 (alphaNCPII), and betaNCP2 (betaNCPII) genes,	
5662	AF230741	complete cds	1,6
5663	AC079878	Homo sapiens BAC clone RP11-343P21 from 7, complete sequence	0.086
		Homo sapiens hypothetical gene supported by AK023961	
5664	XM_029904	(LOC90205), mRNA	1.9
		Arabidopsis thaliana translin-like protein (At2g37020; T1J8.20)	
5665	AY042792	mRNA, complete cds	4.6
5666	AB018347	Homo sapiens mRNA for KIAA0804 protein, partial cds	0.068
		Oryza sativa microsatellite MRG0342 containing (AG)X22, closest to	
5667	AY018017	marker G1185, genomic sequence	0.059
5668	AE005310	Escherichia coli O157:H7 EDL933 genome, contig 1 of 3, section 134 of 155	0.49
		Homo sapiens farnesyl diphosphate synthase (farnesyl pyrophosphate	
		synthetase, dimethylallyltranstransferase, geranyltranstransferase)	
5669	XM_034498	(FDPS), mRNA	5.3
5670	AK027123	Homo sapiens cDNA: FLJ23470 fis, clone HSI11950	0.17
5671	NM_013696	Mus musculus thyrotropin releasing hormone receptor (Trhr), mRNA	0.23
5673	L23758	Bos taurus lysozyme 7A mRNA	0.8
		Arabidopsis thaliana putative polypeptide chain release factor	
5674	AY034963	(F13K23.17) mRNA, complete cds	1.4
5675	XM_028009	Homo sapiens KIAA1563 protein (KIAA1563), mRNA	2E-71
5676	XM_046473	Homo sapiens Rho GDP dissociation inhibitor (GDI) alpha (ARHGDIA), mRNA	0.59
		Homo sapiens fucosyltransferase 1 (galactoside 2-alpha-L-	
5677	NM 000148	fucosyltransferase, Bombay phenotype included) (FUT1), mRNA	0.0000001
5678	AF203477	Drosophila melanogaster Sticky ch1 (stich1) mRNA, partial cds	2.1
		Plasmodium falciparum transmission-blocking target antigen Pfs230	
5679	AF269242	(s230) gene, complete cds	0.23

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	Distant	
NO	ACCESSIN	DESCRIP	P VALUE
5680	AE006954	Mycobacterium tuberculosis CDC1551, section 40 of 280 of the complete genome	2.1
5681	Z93386	Caenorhabditis elegans cosmid R11H6, complete sequence	0.023
5682	XM_012723	Homo sapiens chromosome 18 open reading frame 1 (C18orf1), mRNA	5,8
5683	AF171785	Drosophila melanogaster clone 69 mRNA sequence	0.73
5684	AF159105	Homo sapiens intestinal trefoil factor gene, 5' flanking region	0.02
5685	NM_011889	Mus musculus septin 3 (Sept3), mRNA	4.6
5688	NM 008541	Mus musculus MAD homolog 5 (Drosophila) (Madh5), mRNA	0.0000002
5690	XM 042245	Homo sapiens similar to inner centromere protein antigens (135kD,	
5689	XM_043345	155kD) (H. sapiens) (LOC92175), mRNA	0.075
5000	AT 500100	Human DNA sequence from clone RP11-405J10 on chromosome 10,	
5690	AL590109	complete sequence [Homo sapiens]	0,003
5691	XM 051593	Homo sapiens hypothetical gene supported by AF055004 (LOC93477), mRNA	0.55
5692	U22345	Human chromosome 20q12 locus-specific repeat	0.008
5693	X63385	B.thuringiensis IS231F DNA	0.72
5694	AF288224	Brachyspira hyodysenteriae DNA Gyrase B subunit gene, complete cds	0.006
3094	AF288224	Lycopersicon hirsutum sesquiterpene synthase 2 (SSTLH2) mRNA,	0.006
5695	AF279456	complete cds	0.06
		Homo sapiens Duffy antigen/chemokine receptor (FY) gene, FY*X	
5696	AF055992	allele, complete cds	1.5
		Homo sapiens synuclein alpha interacting protein (SNCAIP) gene,	
5697	AF167306	exon 10 and complete cds	0.99
5698	AJ277985	Nicotiana tabacum drepp1 gene, exons 1-4	0.22
5699	NM_030238	Mus musculus dynein, cytoplasmic, heavy chain I (Dnchc1), mRNA	4E-25
5701	X95343	N.tabacum mRNA for HSR201 protein	5.9
5702	AE002116	Urcaplasma urcalyticum section 17 of 59 of the complete genome	0.68
5704	XM 029907	Homo sapiens anillin (Drosophila Scraps homolog), actin binding protein (ANLN), mRNA	0.72
5705	AJ304804	Homo sapiens ANKRD2 gene for skeletal muscle ankyrin repeat, exons 1-9	
3703	AJ304804	exous 1-9	0.22
5706	AF371373	Cricetulus griseus hypothetical protein A1-3 mRNA, complete cds	0.72
5707	AE007446	Streptococcus pneumoniae section 129 of 194 of the complete genome	0.18
5708	L44121	Homo sapiens (clone pHK2.1D) CMT1A gene, repeat unit	6.6
5709	X90848	S.scrofa ppk98 gene	0.23
5710	AF397903	Pisum sativum AAA-metalloprotease FtsH (FTSH) mRNA, complete cds; nuclear gene for mitochondrial product	3.8
3/10	AF397903	Homo sapiens ADP-ribosylation factor-like 6 interacting protein	3.6
5711	XM_027365	(ARL6IP), mRNA	1E-30

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
5712	L01574	Homo sapiens P-selectin CD62 (GRMP) gene, 5' end and promoter region	0.00001
3/12	L013/4	Oryza sativa microsatellite MRG5943 containing (TTA)X55, closest	0.00001
5713	AY023618	to marker RG908, genomic sequence	0.00009
		Vibrio cholerae chromosome II, section 34 of 93 of the complete	
5714	AE004377	chromosome	0.58
		Botrytis cinerea strain T4 cDNA library under conditions of nitrogen	
5715	AL115472	deprivation	0.2
		Homo sapiens hypothetical gene supported by BC006369; BC008013	
5716	XM_042855	(LOC92105), mRNA	8E-23
		Homo sapiens region containing deleted in azoospermia 2; deleted in	
5717	XM_015203	azoospermia (LOC90902), mRNA	1E-19
		Oryza sativa microsatellite MRG2114 containing (AT)X78, genomic	
5718	AY019789	sequence	0.0000003
5719	AB010300	Garlic virus A genomic RNA, complete sequence	0.001
5720	AF171785	Drosophila melanogaster clone 69 mRNA sequence	0.73
5501	AT225100	Phaseolus coccineus suspensor-specific protein (C541-like) gene,	
5721	AF325188	complete cds	0.24
5722	XM 048692	Homo sapiens similar to hypothetical protein (H. sapiens) (LOC92998), mRNA	1E-16
3122	AIVI_048092	(LOC92998), IIIRINA	1E-10
5723	AE006878	Sulfolobus solfataricus section 237 of 272 of the complete genome	2.2
5724	AF038919	Dictyostelium discoideum PslA (pslA) gene, complete cds	0.026
		Bacillus thuringiensis plasmid encoded Cry1Ba (cry1Ba) gene,	
5725	AF368257	complete cds	2.1
5726	AF148319	Homo sapiens imprinting center, AS-SRO region	1.8
5727	D49412	Human gene for interleukin 3 receptor alpha subunit, exon 11	0.23
5728	AJ012488	Mus musculus gene encoding serotonin receptor 5-HT2B, exons 1-3	0.25
3128	AJ012488	Human DNA sequence from clone RP11-259P20 on chromosome 6.	0.23
5729	AL445224	complete sequence [Homo sapiens]	2
3725	710110221	Homo sapiens cDNA: FLJ21368 fis, clone COL03056, highly similar	_ <u>-</u> _
5730	AK025021	to AF158555 Homo sapiens glutaminase C mRNA	e-171
		Raphanus sativus dark-inducible and senescence-associated gene,	
5731	D38252	exon 1 to 5	0.62
		Homo sapiens hypothetical gene supported by AK026792	
5732	XM_051880	(LOC93533), mRNA	5.6
5733	K00769	yeast (s.pombe) 5s rrna gene and flanks, clone psprl 1	0.023
5734	AL031254	Caenorhabditis elegans cosmid 4R79, complete sequence	0.082
5735	AK001945	Homo sapiens cDNA FLJ11083 fis, clone PLACE1005232	2.1
5736	XM_012452	Homo sapiens ryanodine receptor 3 (RYR3), mRNA	0.008
5737	XM_046442	Homo sapiens hypothetical protein FLJ23045 (FLJ23045), mRNA	6E-33
5738	AF072513	Illex argentinus microsatellite Ia203	0.69
5739	XM_050747	Homo sapiens TC10-like Rho GTPase (TCL), mRNA	5E-20
5740	AC079757	Homo sapiens clone RP11-109N2, complete sequence	0.62

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
5741		Homo sapiens KIAA1691 protein (KIAA1691), mRNA	4.4
5742	NM_004342	Homo sapiens caldesmon 1 (CALD1), mRNA	4E-67
5743	BC002303	Mus musculus, clone IMAGE:3591199, mRNA	0.69
5744	AF273045	Homo sapiens CTCL tumor antigen se14-3 mRNA, complete eds	5E-28
5745	AE001798	Thermotoga maritima section 110 of 136 of the complete genome	0.71
1		Homo sapiens Fanconi anemia complementation group D2 protein	
5746	AF273225	(FANCD2) gene, exon 7	0.009
		Human DNA sequence from clone RP11-117A20 on chromosome 6,	
5747	AL589920	complete sequence [Homo sapiens]	0.026
5748	AF101092	Homo sapiens collagen type XI alpha-1 (COL11A1) genc, exon 29	0.019
5749	S80521	dMax=basic helix-loop-helix/leucine zipper protein {alternatively spliced} [mice, WEHI 231 B-lymphoma cells, mRNA Partial, 752 nt]	0.003
5750	AF291825	Gorilla gorilla von Hippel-Lindau tumor suppressor (VHL) gene, promoter region	0.078
5751	AF120716	Kluyveromyces lactis cytochrome c oxidase subunit III (COX3) gene, mitochondrial gene encoding mitochondrial protein, complete cds	0.027
5752	NM_001423	Homo sapiens epithelial membrane protein 1 (EMP1), mRNA	5E-31
5753	AB045575	Danio rerio DNA, fkk-7c locus, insertion site of the medaka fish Tol2 element	1.7
5754	AL050120	Homo sapiens mRNA; cDNA DKFZp586D211 (from clone DKFZp586D211); partial cds	2
5755	AP000468	Homo sapiens genomic DNA, chromosome 21q22.3, clone:f60A9, complete sequence	0.009
		Homo sapiens cDNA FLJ10787 fis, clone NT2RP4000481, weakly	
5756	AK001649	similar to ATP-DEPENDENT RNA HELICASE DOB1	0
5757	NM 031511	Rattus norvegicus Insulin-like growth factor II (somatomedin A) (Igf2), mRNA	0.025
		Homo sapiens mRNA; cDNA DKFZp434B249 (from clone	
5758	AL162036	DKFZp434B249)	0.71
5760	AF317787	Rattus norvegicus Cd36 gene, promoter region	2.1
5761	XM_032021	Homo sapiens calreticulin (CALR), mRNA	6E-41
5762	XM_037847	Homo sapiens neurexophilin 3 (NXPH3), mRNA	4.2
5763	AF288818	Rattus norvegicus cytochrome P450 4F5 gene, complete cds	0.026
5764	AF269237	Cloning vector pTX-GFP, complete sequence	0.71
5765	AK021493	Homo sapiens cDNA FLJ11431 fis, clone HEMBA1001094	8E-76
5766	AF034077	Equus caballus alpha-1-antitrypsin (Spi2) gene, complete cds	0.0003
5767	XM_030851	Homo sapiens G protein-coupled receptor kinase-interactor 1 (GIT1), mRNA	5.4
5768	AY039919	Arabidopsis thaliana unknown protein (AT4g09340) mRNA, complete cds	4.8

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ACCESSN  AK018211  XM_006622  U72335  XM_048810	Table 3A Nearest Neighbor (BlastN vs. Genbank)  DESCRIP  Mus musculus adult mate medulla oblongata cDNA, RIKEN full- length enriched library, clone:6330510M09, full insert sequence Homo sspicas FTPRF interacting protein, binding protein 1 (liprin beta 1) (PPFIBP1), mRNA  Human platelet activating factor acctylhydrolase, brain isoform, 45 Rba subunit (LSI) gene, exons 3 and 4	P VALUE 6.4 0.079
AK018211 XM_006622 U72335	Mus musculus adult male medulla oblongata cDNA, RIKEN full- length enriched library, clone:6330510M09, full insert sequence Homo sepicns PTPRF interacting protein, binding protein 1 (liprin beta 1) (PPFIBP1), mRNA Human platelet activating factor acetylhydrolase, brain isoform, 45	6.4
AK018211 XM_006622 U72335	Mus musculus adult male medulla oblongata cDNA, RIKEN full- length enriched library, clone:6330510M09, full insert sequence Homo sepicns PTPRF interacting protein, binding protein 1 (liprin beta 1) (PPFIBP1), mRNA Human platelet activating factor acetylhydrolase, brain isoform, 45	6.4
XM_006622 U72335	length enriched library, clone:6330510M09, full insert sequence Homo spicas PTPRF interacting protein, binding protein 1 (liprin beta 1) (PPFIBP1), mRNA Human platelet activating factor acctylhydrolase, brain isoform, 45	
XM_006622 U72335	Homo sapicus PTPRF interacting protein, binding protein 1 (liprin beta 1) (PPFIBP1), mRNA Human platelet activating factor acetylhydrolase, brain isoform, 45	
U72335	beta 1) (PPFIBP1), mRNA Human platelet activating factor acctylhydrolase, brain isoform, 45	0.079
	hDo submit (LIS1) sono suoma 2 and 4	
YM 048810	KDa subunit (LIS1) gene, exons 3 and 4	0.2
YM 048810		
71W_040010	Homo sapiens hypothetical protein FLJ22116 (FLJ22116), mRNA	0
	Arabidopsis thaliana 12-oxo-phytodienoate reductase gene, complete	
AF218257	cds	0.23
AK014450		0.083
'		
AK005342		1.6
		0.063
NC_001619	Canine oral papillomavirus, complete genome	1.8
		0
		0.084
		2.2
XM_040841	Homo sapiens cullin 5 (CUL5), mRNA	0.23
	L	
		0.22
AE006648		0.007
		0.008
		0.23
		1.9
		1.9
		0.69
		2.1
AK001343		2.1
AT190157		6.5
AL 103137		0.5
AK013313		6.1
		0.69
		6E-63
		2E-54
		0.6
237003		- 0.0
AF288409		0.081
1 200400	Compton on	0.001
XM 015806	Homo seniens hypothetical protein EL 192621 (EL 192621) mRNA	l 0
	AK014450  AK005342  M99412 NC_001619 NC_001619 NM_030238 AL031254 274797 XM_040841  U20807 AE006648  AY018666 AY018666 AY018666 AY017281 XM_017281 XM_017281 AF229090 AK0010945 AF189157 AK013313 AK021451 Z34885 AF288408	Mus musculus 18 days pregnant adult female placenta and extra embryonic tissue cDNA, RIKEN füll-length enriched library, clones/330422A13, füll insert sequence  Mus musculus adult male cerebellum cDNA, RIKEN füll-length enriched library, clone:1500032H18, füll insert sequence  My9412 Human interleukin-8 receptor (IL8RB) gene, complete cds  NC_001619 Canine oral papillomavirus, complete genome  NM_030238 Mus musculus dynein, cytoplasmic, heavy chain 1 (Dnchcl.), mRNA AL031254 Caenorhabditis elegans cosmid 4R79, complete sequence  SZ47970 S.cervisiae chromosome XV reading frame ORF YOL055c  XM_040841 Homo supiens cullin 5 (CUL5), mRNA  U20807 Bos taurus protein tyrosine phosphatase BA14 mRNA, complete cds  AE006648 Sulfolobus solitariarius section 7 of 272 of the complete genome  Ovyza sativa microsaelltie MRG0991 containing (AT)X119, closest to  marker R1713, genomic sequence  AY018666 marker R1713, genomic sequence  XM_017281 Homo sepiens cDNA: FLI22646 fis, clone HSI07178  XM_017281 Homo sepiens desulfine alpha (ENSA), mRNA  AF229090 Homo sepiens dendesulfine alpha (ENSA), mRNA  AF229091 Homo sepiens myosin VI (MYO6) gene, exon 11  AK013313 Homo spiens this case virus (strain O1) polyprotein gene, complete  ds  AK013313 Homo spiens filts sease virus (strain O1) polyprotein gene, complete  ds  AK013314 Homo spiens filts gene for serotonin 3 receptor  Homo spiens STRT gene for serotonin 3 receptor  Homo spiens STRT gene for serotonin 3 receptor  Homo spiens STRT gene for serotonin 3 receptor  Homo spiens STRT gene for serotonin 3 receptor  Homo spiens STRT gene for serotonin 3 receptor  Homo spiens STRT gene for serotonin 3 receptor  Homo spiens STRT gene for serotonin 3 receptor  Homo spiens STRT gene for serotonin 3 receptor

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
110	ACCESSIN	Plasmodium falciparum Indochina III/CDC erythrocyte binding	PVALUE
5800	AF332918	protein BAEBL (bacbl) gene, complete cds	3.9
		Homo sapiens hypothetical protein DKFZp547A023	
5801	XM_052063	(DKFZp547A023), mRNA	0
5802	AB013796	Ceratopteris richardii mRNA for CRHB6, complete cds	0.024
		Rattus norvegicus transient receptor potential Trp4beta mRNA,	
5803	AF288408	complete cds	0.082
		Terebra subulata 16S ribosomal RNA gene, partial sequence;	
5804	AF174213	mitochondrial gene for mitochondrial product	2.8
		Homo sapiens serologically defined breast cancer antigen NY-BR-18	
5805	AF308286	mRNA, partial cds	0.59
5806	AF030199	Mus musculus type 1 sigma receptor gene, complete cds	0.49
5807	M99412	Human interleukin-8 receptor (IL8RB) gene, complete cds	0.51
5808	Z34885	T.pyriformis (CGL) gene for TCP1gamma protein	0.58
5809	ATTOREGOO	Mus musculus B lymphocyte induced maturation protein 1 (Prdm1) gene, exon 8 and complete cds	15.65
3609	AF305539	Rattus norvegicus transient receptor potential Trp4beta mRNA,	1E-67
5810	AF288408	complete cds	0.087
5811	AE001181	Borrelia burgdorferi (section 67 of 70) of the complete genome	0.001
3011	12001101	Homo sapiens voltage-dependent calcium channel beta 2 subunit	0.001
5812	AY027894	(CACNB2) gene, exon 2	0.00008
5813	XM 029481	Homo sapiens similar to KIAA0732 protein (H. sapiens) (LOC90154), mRNA	0.68
5814	AK021659	Homo sapiens cDNA FLJ11597 fis, clone HEMBA1003856	2.1
5815	U39721	Mycoplasma genitalium section 43 of 51 of the complete genome	0.19
5816	Z73267	S.cerevisiae chromosome XII reading frame ORF YLR095c	0.009
5817	XM 029453	Homo sapiens oculocerebrorenal syndrome of Lowe (OCRL), mRNA	0.26
		Xenopus laevis kinesin-related protein (XCENP-E) mRNA, complete	
5818	AF027728	cds	0.052
		Plasmodium falciparum chromosome 2, section 44 of 73 of the	
5819	AE001407	complete sequence	0.18
5820	AL157491	Homo sapiens mRNA; cDNA DKFZp434K1111 (from clone DKFZp434K1111)	9E-50
		Homo sapiens cAMP-regulated guanine mucleotide exchange factor II	
5821	XM_002437	(CAMP-GEFII), mRNA	4E-37
5822	XM_050146	Homo sapiens KIAA0472 protein (KIAA0472), mRNA	8E-20
5823	AY032600	Carica papaya xyloglucan endo-transglycosylase mRNA, complete cds	0.19
5824	Z36061	S.cerevisiae chromosome II reading frame ORF YBR192w	3
5825	AK021459	Homo sapiens cDNA FLJ11397 fis, clone HEMBA1000622	0.046
		Nicotiana tabacum putative chloroplast RNA helicase VDL (VDL)	l
		gene, exons 1 through 5 and complete CDS, alternatively spliced;	
5826	AF261017	nuclear gene for chloroplast products	2.1
5827	AJ277162	Beta vulgaris mRNA for putative Cdc2-related protein kinase CRK2	0.009

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
1 1		Human DNA sequence from clone RP11-238K16 on chromosome 9,	
5828	AL390874	complete sequence [Homo sapiens]	0.026
		Solanum tuberosum type A phytochrome (c-phyA) mRNA, complete	
5829	S84872	cds	0.43
		Homo sapiens interphotoreceptor matrix proteoglycan 200 (IMPG2)	
5830	AF271370	gene, exon 10	0.078
5831	XM_051183	Homo sapiens embryonic ectoderm development (EED), mRNA	0.00002
		Homo sapiens, serum/glucocorticoid regulated kinase 2, clone	
5832	BC006523	IMAGE:2988475, mRNA	3E-34
5833	AF100658	Caenorhabditis elegans cosmid H08G01	0.44
5834	XM_003725	Homo sapiens putative ribonuclease III (RNASE3L), mRNA	0.008
		Amoeba proteus symbiotic bacterium 29kDa protein (s29x) gene,	
5835	M58029	complete cds_	0.71
5836	U46541	Staphylococcus aureus sarA gene, complete cds	0,003
5837	AJ225333	Acidianus ambivalens plasmid pDL10	0.17
5838	X15702	Fruitfly LA9 mRNA for DNA binding protein	0.077
		Human DNA sequence from clone CTA-232D4 on chromosome	
5839	AL031590	22q13.1 Contains a GSS, complete sequence [Homo sapiens]	0.16
5840	X07946	Yeast plasmid DNA coding for RNA polymerase subunit	0.082
		Human DNA sequence from clone RP11-479D11 on chromosome 10,	
5841	AL590324	complete sequence [Homo sapiens]	0.082
5842	XM_004947	Homo sapiens paraoxonase 2 (PON2), mRNA	1.6
		B 114 - 41 - 4 - 4 - 4 - 4 - 4 - 4 - 4 -	
5843	AF262582	Dolichorhinotermes sp. 'Manaus' 16S mitochondrial ribosomal RNA gene, partial sequence; mitochondrial gene for mitochondrial product	0,067
3843	AF202362	Homo sapiens (subclone 2 h3 from P1 H43) DNA sequence, complete	0,007
5844	L81839	sequence   Subcrone 2_113 from P1 H43) DNA sequence, complete	0,00006
3044	F01033	sequence	0.00000
5845	U67550	Methanococcus jannaschii section 92 of 150 of the complete genome	0.087
3013	007330	Homo sapiens hypothetical gene supported by U00951 (LOC92415).	0.007
5846	XM 044987	mRNA	6.1
3010	2047307	Plasmodium falciparum chromosome 2, section 57 of 73 of the	0.1
5847	AE001420	complete sequence	0.71
5011	110001120	Stylonychia mytilus DNA-dependent RNA polymerase II largest	
5848	AF315823	subunit RPB1 (RPB1) gene, partial cds	0.51
		Physcomitrella patens chloroplast genes for ribulose-1,5-bisphosphate	
		carboxylase/oxygenase large subunit, tRNA-Arg and acetyl-CoA	
5849	AB066207	carboxylase beta subunit, complete cds	0.65
		Plasmodium chabaudi adami carbamoyl phosphate synthetase II gene,	
5850	AF286898	partial cds	0.002
		Human DNA sequence from clone RP11-132E18 on chromosome 10,	
5851	AL583888	complete sequence [Homo sapiens]	0.028
		Chromatium vinosum genes for light-harvesting and reaction center	
5852	AB011811	proteins, partial and complete cds	6.4
5853	AF153448	Zea mays nitrate reductase (NR1) gene, complete cds	0.067

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	<u> </u>	Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
5854	XM 012762	Homo sapiens hypothetical protein FLJ10891 (FLJ10891), mRNA	0
ì		Maoricicada cassiope isolate TB-MI-004 (RNA-Asp gene, complete	)
		sequence; ATPase subunit 8 gene, complete cds; and ATPase subunit	
5855	AF248814	6 gene, partial cds; mitochondrial genes for mitochondrial products	0.029
		Anthocidaris crassispina mRNA for dyncin beta-heavy chain,	
5856	D01021	complete cds	2E-22
		Plasmodium falciparum chromosome 2, section 38 of 73 of the	
5857	AE001401	complete sequence	0.54
5858	BC007548	Homo sapiens, clone IMAGE:2959994, mRNA	1.9
		Macaca fascicularis brain cDNA clone:QmoA-12555, full insert	
5859	AB063065	sequence	0.18
		Mus musculus myristoylated alanine rich protein kinase C substrate	
5860	NM_008538	(Macs), mRNA	1.5
5861		Homo sapiens sarcoglycan, epsilon (SGCE), mRNA	0.66
5862	AE006685	Sulfolobus solfataricus section 44 of 272 of the complete genome	0.37
5863	BC008007	Homo sapiens, clone IMAGE:3510538, mRNA	6E-34
		Nicotiana tabacum mRNA for TMV response-related gene product,	
5864	AB024511	complete cds	2.1
		Glycine max phosphatidylinositol-specific phospholipase C mRNA,	
5865	U25027	complete cds	0.077
5866	U50715	Mus musculus alpha-galactosidase A gene, complete cds	0.024
		Stylophora pistillata L-type calcium channel alpha-1 subunit	
5867	U64465	(STPCACHL) mRNA, complete cds	0.072
1		Homo sapiens similar to hypothetical protein DKFZp566D1346 (H.	}
5868	XM_040043	sapiens) (LOC91690), mRNA	0.023
Į l		Arabidopsis thaliana serine/threonine protein phosphatase 2A-3	Į.
5869	U60135	catalytic subunit gene, complete cds	0.027
ì		Lycopersicon esculentum plasma membrane H+-ATPase (LHA2)	1
5870	AF275745	mRNA, complete cds	1.5
		C.crispus Chloroplast gene encoding 16S rRNA, tRNA-Ile, tRNA-	
5871	Z29521	Ala, and 23S rRNA (partial)	0.003
5872	AJ132557	Oryctolagus cuniculus CYP19 gene, ovarian promoter region	0.24
5873	AF124524	Arabidopsis thaliana gamma-adaptin 1 gene, complete cds	0.24
·		Mus musculus 10, 11 days embryo cDNA, RIKEN full-length	
5874	AK013131	enriched library, clone:2810422B04, full insert sequence	0.72
5875	AB044413	Sus scrofa MMP-3 mRNA for matrix metalloproteinase-3, partial cds	0.23
		Porcine adenovirus 3 171R (E1A), 202R (E1B-1), 474R (E1B-2),	
1		288R (288R), 198R (pIX), 163R* (163R*), 162R (162R), 97R (97R)	1
		and 184R (184R) genes, complete cds; and IVa2 (IVa2) gene, partial	
5876	AF247039	cds	5.2
5877	AF060579	Gossypium barbadense clone pXP020 repetitive DNA sequence	0.055
		Streptococcus pyogenes M1 GAS strain SF370, section 28 of 167 of	
5878	AE006499	the complete genome	2.1

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID NO	ACCESSN	DESCRIP	P VALUE
5879	AL591498	Human DNA sequence from clone RP11-113L12 on chromosome 13, complete sequence [Homo sapiens]	0,003
5880	NM_017066	Rattus norvegicus Pleiotrophin (Heparine binding factor, Hbnf, in the mouse) (Ptn), mRNA	0.21
5881	XM_047505	Homo sapiens hypothetical gene supported by AK022483 (LOC92818), mRNA	0.7
5882	J00912	chicken brain tubulin, alpha chain mrna	0.24
5883	M15840	Human interleukin 1-beta (IL1B) gene, complete cds	0,08
5884	X92683	H. sapiens DNase I hypersensitive site (HSS-1)	2.1
		Human DNA sequence from clone RP1-68P15 on chromosome 11p13- 14.2 Contains GSSs and ESTs. Contains part of a novel gene.	
5885	AL390767		1.0
5886	XM 012386	complete sequence [Homo sapiens]	1.8 1E-48
2000	AM_012380	Homo sapiens KIAA0377 gene product (KIAA0377), mRNA	1E-48
5887	AL137466	Homo sapiens mRNA; cDNA DKFZp434H1322 (from clone DKFZp434H1322)	0.00001
5888	X62745	Z.mays OBF1 mRNA for ocs-element binding factor 1	0.23
5890	AB048902	Macaca fascicularis brain cDNA, clone:QnpA-17448	0.24
5892	AF221070	Ophiostoma piliferum strain CBS129.32 18S ribosomal RNA gene, partial sequence; internal transcribed spacer 1, 5.8S ribosomal RNA gene, and internal transcribed spacer 2, complete sequence	0,003
5893	AE006530	Streptococcus pyogenes MI GAS strain SF370, section 59 of 167 of the complete genome	0,22
5894	AB018698	Broad bean wilt virus 2 gene for precursor polyprotein, complete cds, isolate: IP	0.025
5895	U67537	Methanococcus jannaschii section 79 of 150 of the complete genome	0.026
5896	AJ000742	Homo Sapiens hisH1 gene, 5' UTR	0,008
5897	AF247039	Porcine adenovirus 3 171R (E1A), 202R (E1B-1), 474R (E1B-2), 288R (288R), 198R (pIX), 163R* (163R*), 162R (162R), 97R (97R) and 184R (184R) genes, complete cds; and IVa2 (IVa2) gene, partial cds	5.2
5898	AJ290290	Asphodeline lutea chloroplast trnL-trnF intergenic spacer	0.23
5899	XM_042440	Homo sapiens UDP-Gal:betaGlcNAc beta 1,4- galactosyltransferase, polypeptide 3 (B4GALT3), mRNA	6.4
5900	U78090	Rattus norvegicus potassium channel regulator 1 mRNA, complete cds	2.1
5901	NM_017053	Rattus norvegicus Neuromedin K receptor (Neurokinin B/Tachikin 3) (Tac3r), mRNA	1.9
5902	AF208658	Tapesia yallundae eburicol 14 alpha-demethylase (CYP51) gene, complete cds	6,2
5903	AF281786	HIV-1 isolate DK-g3 from USA gag polyprotein (gag) gene, partial cds	0.63
5904	AJ011019	Capra hircus csn2 gene, exons 1 to 9, allele 0(null)	0.63
3304	7110119	Chironomus pallidivittatus 4L ORF for putative recombinase	
5905	AJ311054	subtelocentric clone, Cp5.5	0.023

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
5906	Y17466	Fugu rubripes SIAT3C gene, last three exons	0.65
5907	AK022127	Homo sapiens cDNA FLJ12065 fis, clone HEMBB1002249	0.001
5908	X79100	C.vicina (B11) mRNA for arylphorin receptor	0.6
5909	AK023079	Homo sapiens cDNA FLJ13017 fis, clone NT2RP3000628	0.23
		Mus musculus 18 days pregnant adult female placenta and extra	
		embryonic tissue cDNA, RIKEN full-length enriched library,	
5910	AK014448	clone;3830421J05, full insert sequence	1.4
		Mus musculus adult male cecum cDNA, RIKEN full-length enriched	
5911	AK020293	library, clone:9130404H11, full insert sequence	0.23
		Lactococcus lactis subsp. lactis IL1403 section 47 of 218 of the	
5912	AE006285	complete genome	0.25
5012	TT41000	Harmon and ania (DDELD) and SLG-1-in-	0.001
5913	U41292	Human prolargin (PRELP) gene, 5' flanking sequence and exon 1	0,001
5914	AL109707	Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 295344	0.00004
3914	AL109/07	Rattus norvegicus proton gated cation channel ASIC1 (Accn2).	0.00004
5915	NM 024154	mRNA	6.2
3713	144 024154	Plasmodium berghei extrachromosomal plastid PB-2, tRNA-Pro,	0.2
		tRNA-Glu, tRNA-Lys, tRNA-Asp, tRNA-Ser, tRNA-Tyr, tRNA-Met,	
ł		tRNA-Leu, tRNA-Cys, and tRNA-His genes, complete sequence, rps4	
		gene, complete cds, tRNA-Thr gene, complete sequence, and large	
5916	U79732	subu>	0.023
		Coniglobus mercatorius 16S ribosomal RNA gene, mitochondrial	
5917	AF098715	gene encoding mitochondrial product, partial sequence	0.095
		Toxoplasma gondii ycf24 protein (ycf24) gene, partial cds; DNA	
	\	dependent RNA polymerase beta subunit (rpoB) gene, complete cds;	
İ		and DNA dependent RNA polymerase beta' subunit (rpoC1) gene,	
5918	AF095904	plastid genes encoding plastid proteins, partial cds	0.026
5919	XM_027416	Homo sapiens mesothelin (MSLN), mRNA	0.7
		Mus musculus, Similar to inhibitor of kappa light polypeptide gene	
5920	BC004772	enhancer in B-cells, kinase beta, clone IMAGE:3256890, mRNA	1.6
5921	AK000935	Homo sapiens cDNA FLJ10073 fis, clone HEMBA1001731	0.18
		Homo sapiens, clone MGC:12595 IMAGE:4303422, mRNA,	
5922	BC009942	complete cds	0.001
5923	J05008	Homo sapiens endothelin-1 (EDN1) gene, complete cds	0.5
		Chlamydophila pneumoniae AR39, section 10 of 94 of the complete	
5924	AE002174	genome	0.69
5925	AJ223616	Caehorhabditis elegans mRNA for calcium ATPase	0.7
		Dictyostelium discoideum prespore-specific protein (pspC) gene,	
5926	AF310894	partial cds; RacH (racH) gene, complete cds; and BOP (bopA) gene,	0.027
3920	AF310894	partial cds	0.027
5927	AE003910	Xylella fastidiosa 9a5c, section 56 of 229 of the complete genome	0.73
3321	AL003910	Ayrona rasumosa 9a.5c, section 50 of 229 of the complete genome	0.73
5928	AF335424	Homo sapiens sperm protein Sp17-2 pseudogene, complete sequence	1.8

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID		The six Indicat Perguoti (Diastri vs. Gellodik)	
NO NO	ACCESSN	DESCRIP	P VALUE
5929	XM_016924	Homo sapiens hypothetical protein MGC4054 (MGC4054), mRNA	4.3
		Mycoplasma capricolum NADH oxidase (naox) gene, partial cds, and	
		lipoate-protein ligase (lpla), pyruvate dehydrogenase EI alpha subunit	
		(odpa), pyruvate dehydrogenase EI beta subunit (odpb), pyruvate	
5930	U62057	dehydrogenase EII (odp2), dihydrolipoamide dehydroge>	0.72
		Homo sapiens DNA methyltransferase (DNMT1) gene, exons 2, 3,	
5931	AF202552	and 4	2.2
5932	AF010604	Homo sapiens SMAD5 (Smad5) gene, exon 3	0.0001
5933	AC084157	Caenorhabditis elegans cosmid Y46E12BR, complete sequence	0.71
		Homo sapiens hypothetical protein DKFZp564D172	
5934	XM_027907	(DKFZP564D172), mRNA	0.18
		Arabidopsis thaliana unknown protein (T9C5.150/AT3g49560)	
5935	AF360191	mRNA, complete cds	6.3
		Culex pipiens Twin-Cp5 SINE retroposon, genomic sequence and v-	
5938	AF282728	SNARE-like gene, partial sequence	0.066
5939	M81702	Candida boidinii methanol oxidase (AOD1) gene, complete cds	0.008
		Homo sapiens 17-beta-hydroxysteroid dehydrogenase IV (HSD17B4)	
5940	AF057740	gene, exon 24 and complete cds	0.21
5941	XM_031463	Homo sapiens exonuclease NEF-sp (LOC81691), mRNA	0.072
5942	AB022048	Mus musculus gene for prolyl oligopeptidase, exon 3, 4, 5, 6	2.1
		R.violacea chloroplast genes for alpha and beta subunit of	
5943	Z48165	phycocyanin	2.1
5944	AF181251	Detter and the Name of the Control o	
3944	AF181251	Rattus norvegicus lung Kruppel-like factor (Lklf) gene, complete cds Homo sapiens genomic DNA, chromosome 21, clone:D40G11, MX1-	5.9
5945	AP001057	D21S171 region, complete sequence	0.00001
3943	AF001037	Mus musculus adult male testis cDNA. RIKEN full-length enriched	0.00001
5946	AK016374	library, clone:4930589L23, full insert sequence	5.9
5947	M27697	Pig urate oxidase mRNA, complete cds	2.1
3747	1012/09/	Fig thate oxidase mixtox, complete cus	2.1
		Homo sapiens similar to DESTRIN (ACTIN DEPOLYMERIZING	
5948	XM 032603	FACTOR) (ADF) (H. sapiens) (LOC90563), mRNA	0.69
5949	U39688	Mycoplasma genitalium section 10 of 51 of the complete genome	0.077
3747	037088	iviyoopiasiia geimanum seetion 10 ot 51 or the complete genome	0.077
		Lactobacillus delbrueckii YmdA (ymdA) gene, partial cds; and	
		putative undecaprenyl-phosphate N-acetyl-glucosaminyl transferase	
		(rgpG), YvyE (yvyE), putative ComF1 protein (comF1), putative	1
5950	AF320250	ComF3 protein (comF3), YvyD (yvyD), putative preprotein translo>	0.69
5951	Y09852	H.sapiens FGFR3 gene, partial	2E-10
		Property of the Board, Property	
5952	Z79103	H.sapiens flow-sorted chromosome 6 HindIII fragment, SC6pA22F11	0.00002
		Homo sapiens genomic DNA, chromosome 21q22.1, D21S226-AML	7
5953	AP000321	region, clone:Q82F5, complete sequence	0.00004
5954	AK025042	Homo sapiens cDNA: FLJ21389 fis, clone COL03455	2E-66

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	-
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
5955	U14576	Dictyostelium discoideum filopodin (talA) gene, complete cds	0.21
		Plasmodium falciparum chromosome 2, section 42 of 73 of the	
5956	AE001405	complete sequence	0.052
5957	M64861	D.discoideum TFIID mRNA, complete cds	0.002
5958	AJ245435	Plasmodium falciparum gc-alpha gene for guanylyl cyclase alpha	0.0001
5959	XM_031443	Homo sapiens phosphodiesterase 8A (PDE8A), mRNA	8E-13
5960	AC079875	Homo sapiens clone CTD-2333B13, complete sequence	0.00003
		Schizophyllum commune unknown gene; mitochondrial gene for	
5961	AF275272	mitochondrial product	0.44
		H.sapiens CpG island DNA genomic Mse1 fragment, clone 18e1,	
5962	Z60119	reverse read cpg18e1.rt1b	0.008
5963	AF100654	Caenorhabditis elegans cosmid C24E9	0.026
		Plasmodium falciparum chromosome 2, section 36 of 73 of the	
5964	AE001399	complete sequence	0.001
5965	U89019	Hepatitis C virus polyprotein gene, complete cds	0.2
		Homo sapiens ARP3 (actin-related protein 3, yeast) homolog	
5966	XM_052512	(ACTR3), mRNA	2E-32
	*****	Human chromosome 21, Down syndrome critical region transcript,	
5967	U17634	SP6 end of clone 3-3-f8	0.7
5968	AF197477	Homo sapiens TRBP pseudogene, partial sequence	0.22
5050	*****	Streptococcus mutans putative D,D-carboxypeptidase and putative N-	
5969	U78599	acetyl-muramidase genes, partial cds	2.1
5970	4.D001414	Homo sapiens genomic DNA, chromosome 21q22.2, clone:D30,	
3970	AP001414	LB7T-ERG region, complete sequence	2.1
5971	AJ277067	Homo sapiens partial SSH3BP1/e3B1 gene for spectrin SH3 binding protein, exon 3-5	5.7
5972	X13294	Human mRNA for myb-related gene A-myb 5'-region	5.7 6E-74
5973	NC 001796	Human parainfluenza virus 3, complete genome	0.053
3713	NC_001790	Hordeum vulgare mRNA for alpha-keto acid dehydrogenase-like	0.053
5974	AJ222787	protein, clone RG136	0,073
37,14	AJZZZ/07	Melanogrammus aeglefinus vitellogenin B (VtgB) mRNA, complete	0.073
5975	AF284034	cds	2.1
3773	A1 204034	Arabidopsis thaliana ubiquitin-protein ligase 2 (UPL2) gene, complete	2.1
5976	AF127565	cds	4,3
3770	211 12/303	Homo sapiens genomic DNA, chromosome 21q22.2, clone:PAC24K9,	4,3
5977	AP001419	LB7T-ERG region, complete sequence	0,0002
3711	711 001417	Human DNA sequence from clone RP11-349A16 on chromosome	0,0002
5978	AL590384	Xq22.3-24, complete sequence [Homo sapiens]	0.02
3776	7000004	Homo sapiens TSC2, NTHLI/NTH1 and SLC9A3R2/E3KARP genes,	0.02
5979	AB014460	partial and complete cds	0,69
05/5	112011100	partial and complete cus	0.03
5980	YM 018334	Homo sapiens hypothetical protein FLJ22418 (FLJ22418), mRNA	0.026
5981	AF229850	Arabidopsis thaliana ferritin (Fer1) gene, complete cds	0.69
	. 11 227 030	Mus musculus 10 days neonate head cDNA, RIKEN full-length	0.07
5982	AK017445	enriched library, clone:5530402H04, full insert sequence	2.1
5983	AF197159	Mus musculus cubilin mRNA, partial cds	0.7
لتنت		The state of the s	

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		Table 3A Nearest Neighbor (BlastN vs. Genbank)	
SEQ ID			
NO	ACCESSN	DESCRIP	P VALUE
		Homo sapiens hypothetical gene supported by U79302; AK026607	
5984	XM_029346	(LOC90138), mRNA	0.026
5985	AK021517	Homo sapiens cDNA FLJ11455 fis, clone HEMBA1001497	0.05
5986	Z74823	S.cerevisiae chromosome XV reading frame ORF YOL081w	0.23
		Homo sapiens KIAA0001 gene product; putative G-protein-coupled	
		receptor; G protein coupled receptor for UDP-glucose (KIAA0001),	}
5987	XM_003091	mRNA	1.9
		Homo sapiens similar to NONHISTONE CHROMOSOMAL	
5988	XM_039450	PROTEIN HMG-14 (H. sapiens) (LOC91606), mRNA	4E-47
		Homo sapiens G protein coupled receptor interacting protein,	
5989	XM_045103	complement-c1q tumor necrosis factor-related (ZSIG37), mRNA	0.23
5990	AK026086	Homo sapiens cDNA: FLJ22433 fis, clone HRC09172	0.078
5991	Z22800	H.sapiens microsatellite repeat	3E-45
		Mycoplasma mycoides mycoides LC hypothetical surface located	
		membrane protein and lipoprotein B precursor (lppB) genes, partial	
5992	AF165136	cds	0.23
		Staphylococcus epidermidis strain SR1 clone step.1004h05 genomic	
5993	AF269465	sequence	0.23
5994	XM_049474	Homo sapiens KIAA1538 protein (KIAA1538), mRNA	0.23
		Homo sapiens (subclone 1_h7 from BAC H112) DNA sequence,	
5995	AC001478	complete sequence	0.63
		Influenza A virus (A/Goose/Guangdong/3/97(H5N1)) segment 3	
5996	AF380163	polymerase (PA) gene, complete cds	0.077
5997	Z74263	S.cerevisiae chromosome IV reading frame ORF YDL215c	0.36
5998	AF282059	Homo sapiens clone 2qtel_edc17fR sequence	7.9
5999	AF233898	Escherichia coli strain KI1218 BfpA (bfpA) gene, partial cds	0.073
6000	M92086	Wasp venom protein mRNA sequence	1.7
6001	XM_006601	Homo sapiens hypothetical protein FLJ10659 (FLJ10659), mRNA	0
6002	D14074	Bovine mRNA for cyclophilin, complete cds	0.23
6003	AC079391	Homo sapiens clone CTD-2243M11, complete sequence	4
6004	AE006055	Pasteurella multocida PM70 section 22 of 204 of the complete genome	0.077
6005	Y09000	R.norvegicus mRNA for dendrin	0.18
		Mus musculus adult male testis cDNA, RIKEN full-length enriched	
6006	AK006991	library, clone:1700084F23, full insert sequence	2.1
		Pelargonium peltatum cultivar Guenievre microsatellite Pp3a	
6007	AF319636	sequence	0.001
6008	D17668	Clostridium septicum gene for alpha-toxin, complete cds	0.077
6009	AK025193	Homo sapiens cDNA: FLJ21540 fis, clone COL06156	0.025
6010	X17191	E.gracilis chloroplast RNA polymerase rpoB-rpoC1-rpoC2 operon	0.22

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		T TORGETS)
NO	N	DESCRIPTION	P VALUE
		gi 12721131 gb AAK02908.1  (AE006121) unknown	
4	12721131	[Pasteurella multocida]	0.14
	ł	gi 1504016 dbi BAA13207.1  (D86971) no similarities	
6	1504016	to reported gene products [Homo sapiens]	0.0003
		gi 12838708 dbj BAB24303.1  (AK005889) putative	
7	12838708	[Mus musculus]	5E-74
		gi 12854246 dbj BAB29972.1  (AK015778) putative	
8	12854246	[Mus musculus]	2.2
		gi 14715574 dbj BAB62041.1  (AB065434) PRPK	
9	14715574	mutant1 [Homo sapiens]	4E-80
		gi 6678964 rcf NP 032671.1  murinoglobulin 1 [Mus	
		musculus] sp P28665 A2M1 MOUSE	
		MURINOGLOBULIN 1 PRECURSOR (MUG1)	
		gb AAA73048.1  (M65736) [Mouse murinoglobulin	
10	6678964	mRNA, complete cds.], gene product [Mus musculus]	4.2
		, , , , , , , , , , , , , , , , , , , ,	
	Ì	gi 11351090 pir  E82999 probable beta-ketoacyl	
		synthase PA5174 [imported] - Pseudomonas acruginosa	
ĺ		(strain PAO1) gb AAG08559.1 AE004930 5	
		(AE004930) probable beta-ketoacyl synthase	
14	11351090	[Pseudomonas aeruginosa]	9.1
		[- ************************************	
		-1050 CTC11 - (\$100 0 C2222 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	
		gi 9506761 ref NP_062298.1  glucosamine-phosphate N-	
	1	acetyltransferase; glucosamine-6-phosphate	
		acetyltransferase [Mus musculus] emb CAA04463.1	
		(AJ001006) EMeg32 protein [Mus musculus] dbj BAB22120.1  (AK002466) putative [Mus musculus]	
1	1	dbj BAB22120.1  (AK002466) putative [Mus musculus]   dbj BAB25161.1  (AK007647) putative [Mus musculus]	
		dbj BAB2512.1  (AK007647) putative [Mus musculus]   dbj BAB25212.1  (AK007722) putative [Mus musculus]	
		dbj[BAB25240.1] (AK007764) putative [Mus musculus]	
		dbj BAB25749.1  (AK008566) putative [Mus musculus]	
		dbj BAB27395.1  (AK0108300) putative [Mus musculus]	
15	9506761	dbj[BAB31241.1] (AK018499) putative [Mus musculus]	1E-84
-13	2200701	gi 9369403 gb AAF87151.1 AC002423 16 (AC002423)	112-04
20	9369403	T23E23.19 [Arabidopsis thaliana]	7.4
-20	2302403	gi 12862430 dbj BAB32466.1  (AB047280) Gag-like	7.4
24	12862430	protein [Tricholoma matsutake]	2.6
	12002-30	Proton [ 1110101011111 Handulako]	2.0

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
l		gi 7434102 pir  H71620 aspartate transaminase (EC	
l		2.6.1.1) PFB0200c [similarity] - malaria parasite	
2.5	7434102	(Plasmodium falciparum) gb AAC71829.1  (AE001380) aspartate aminotransferase [Plasmodium falciparum]	0.72
23	7434102	gi 14721245 ref XP 044013.1  kinesin-like protein 2	0.73
26	14721245	[Homo sapiens]	8E-91
-20	14721243	[Попо заролз]	0E-91
		Norocettal (DTD octored)	
		gi 9506713 ref[NP_061856.1  nucleolar protein family	
		A, member 1; H/ACA small nucleolar RNPs protein 1 [Homo sapiens] ref[XP 003601.1  nucleolar protein	
		family A, member 1 (H/ACA small nucleolar RNPs)	
		[Homo sapiens] ref[XP 039016.1] nucleolar protein	
ĺ		family A, member 1 (H/ACA small nucleolar RNPs)	
		[Homo sapiens] ref[NP_127460.1  nucleolar protein	
		family A, member 1; H/ACA small nucleolar RNPs	
ł		protein 1 [Homo sapiens] emb CAB76563.1	
		(AJ276003) GAR1 protein [Homo sapiens]	
		gb AAH03413.1 AAH03413 (BC003413) nucleolar	
		protein family A, member 1 (H/ACA small nucleolar	
27	9506713	RNPs) [Homo sapiens]	4
		gi 14150082 ref NP_115691.1  hypothetical protein	
		MGC4399 [Homo sapiens] gb AAH04991.1 AAH04991 (BC004991) Similar to RIKEN cDNA 5730438N18	
28	14150082	gene [Homo sapiens]	0.001
	1110000	gi 12861366 dbj BAB32182.1  (AK020701) putative	- 0.001
29	12861366	[Mus musculus]	5E-42
32	14739173	gi 14739173 ref XP_037531.1  59395 [Homo sapiens]	3E-23
		gi 7513572 pir  S72438 phosphatidylserine	
		decarboxylase (EC 4.1.1.65) precursor, mitochondrial -	
36	7513572	Chinese hamster	e-112
		gi 7488381 pir  T08408 transcription factor homolog	
		F18B3.150 - Arabidopsis thaliana emb CAB42916.1	
27	7488381	(AL049862) transcription factor-like protein [Arabidopsis thaliana]	0.1
37	/488381	[Arabidopsis thaliana] gi 12849752 dbi BAB28466.1  (AK012782) putative	8.1
38	12849752	[Mus musculus]	5E-89
<del></del>	-20.5.52	gi 14715574 dbi BAB62041.1  (AB065434) PRPK	3207
39	14715574	mutant1 [Homo sapiens]	1E-88
		gi 12832845 dbj BAB22281.1  (AK002682) putative	
40	12832845	[Mus musculus]	4E-51

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 12832845 dbj BAB22281.1  (AK002682) putative	
43	12832845	[Mus musculus]	6E-45
		gi 11558040 cmb CAC17800.1  (AJ245890)	
47	11558040	hypothetical protein [Polymyxa betae]	7.6
		gi 8052306 emb CAB92239.1  (AJ242721) AIP37	
48	8052306	protein [Mus musculus]	1.3
		gi 13812380 ref NP_113498.1  N-myristoyltransferase	
		[Guillardia theta] emb CAC27067.1  (AJ010592) N-	100
49	13812380	myristoyltransferase [Guillardia theta]	7
		gi 8885549 dbj BAA97479.1  (AB025604) telomere	
51	8885549	repeat-binding protein [Arabidopsis thaliana]	1.5
		gi 14727244 ref XP 035596.1  hypothetical protein	
		AF140225 [Homo sapiens] ref[XP_035597.1] 49482	
		[Homo sapiens] gb AAH07829.1 AAH07829	
		(BC007829) Similar to hypothetical protein AF140225	
52	14727244	[Homo sapiens]	6E-58
		gi 14737646 ref XP_038048.1  similar to APICAL	
		ENDOSOMAL GLYCOPROTEIN PRECURSOR (R.	
57	14737646	norvegicus) [Homo sapiens]	0.003
		gi 13811699 ref NP_109668.1  zinc finger protein 313;	
		zinc finger protein 228 [Mus musculus]	
		gb AAG01141.1 AF282919_1 (AF282919) Zfp228	
58	13811699	[Mus musculus]	0.00002
		gi 14042394 dbj BAB55226.1  (AK027604) unnamed	
59	14042394	protein product [Homo sapiens]	e-112
60	14769036		9E-97
		gi 13648339 ref XP_003184.2  zinc finger protein	
61	13648339	ANC_2H01 [Homo sapiens]	0.0006
		gi 13638702 ref XP_015971.1  hypothetical protein	
		FLJ23384 [Homo sapiens] ref[XP_048858.1]	
64	13638702	hypothetical protein FLJ23384 [Homo sapiens]	2E-86
		gi 14150082 ref NP_115691.1  hypothetical protein	
		MGC4399 [Homo sapiens] gb AAH04991.1 AAH04991	
		(BC004991) Similar to RIKEN cDNA 5730438N18	
65	14150082	gene [Homo sapiens]	0.00003
		gi 14727244 ref XP_035596.1  hypothetical protein	
		AF140225 [Homo sapiens] ref[XP_035597.1] 49482	
		[Homo sapiens] gb AAH07829.1 AAH07829	
		(BC007829) Similar to hypothetical protein AF140225	
67	14727244	[Homo sapiens]	2E-38

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	Table 3B Treatest (reignoof (Blasta Vs. Non-Redundam	riotells)
NO	N	DESCRIPTION	P VALUE
		gi 14727244 ref XP 035596.1  hypothetical protein	
		AF140225 [Homo sapiens] ref[XP 035597.1] 49482	
		[Homo sapicns] gb AAH07829.1 AAH07829	
		(BC007829) Similar to hypothetical protein AF140225	
68	14727244	[Homo sapiens]	6E-60
		gi 14715574 dbj BAB62041.1  (AB065434) PRPK	
69	14715574	mutant1 [Homo sapiens]	2E-98
		gi 7491888 pir  T41272 hypothetical protein	
		SPCC297.05 - fission yeast (Schizosaccharomyces	
]		pombe) emb CAB40785.1  (AL049609) hypothetical	
70	7491888	protein [Schizosaccharomyces pombe]	9.3
		gi 12843000 dbj BAB25820.1  (AK008666) putative	
71	12843000	[Mus musculus]	1E-86
		gi 13542310 ref[NP_111998.1  Uncharacterized	
72	13542310	conserved protein [Thermoplasma volcanium]	6.6
		gi 14728906 ref XP_003327.3  hypothetical protein	
73	14728906	FLJ10858 [Homo sapiens]	7E-98
	,	gi 14749106 ref XP_031512.1  14599 [Homo sapiens]	
		dbj BAB15298.1  (AK025963) unnamed protein product	
74	14749106	[Homo sapiens]	1E-26
		gil1176169lsplP28636lYHBE ECOLI	
İ		HYPOTHETICAL 35.0 KDA PROTEIN IN DACB-	
		RPMA INTERGENIC REGION pir B65109	
		hypothetical 35.0K protein (dacB-rpmA intergenic	
		region) - Escherichia coli gb AAA57985.1  (U18997)	
		ORF f321 [Escherichia coli] gb AAC76216.1	
		(AE000399) orf, hypothetical protein [Escherichia coli	
		K12] gb AAG58318.1 AE005547_4 (AE005547) orf,	
		hypothetical protein [Escherichia coli O157:H7	
		EDL933] dbj BAB37486.1  (AP002564) hypothetical	
76	1176169	protein [Escherichia coli O157:H7]	0.23
		gi 13161357 dbj BAB32948.1  (AP002908) hypothetical	
		protein~similar to Arabidopsis thaliana F1P2.140	
77	13161357	[Oryza sativa]	6.3
		gi 8922806 ref NP_060761.1  hypothetical protein	
		FLJ10986 [Homo sapiens] dbj BAA91940.1	
		(AK001848) unnamed protein product [Homo sapiens]	
		gb AAH00610.1 AAH00610 (BC000610) hypothetical	
78	8922806	protein FLJ10986 [Homo sapiens]	5E-83
		gi 14603356 gb AAH10136.1 AAH10136 (BC010136)	
82	14603356	Unknown (protein for MGC:19513) [Homo sapiens]	1E-74

	Ι	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	Transfer (Stability 10: 1101 100 1101	1 Totoms/
NO	N	DESCRIPTION	P VALUE
		gi 12804667 gb AAH01758.1 AAH01758 (BC001758)	
83	12804667	hypothetical protein FLJ20758 [Homo sapiens]	0.00000005
	Į.		
00	0705600	gi 9795608 gb AAF98426.1 AC021044_5 (AC021044)	
90	9795608	Unknown protein [Arabidopsis thaliana]	0.7
91	12621027	gi 13631937 ref XP_001412.3  mctal-regulatory transcription factor 1 [Homo sapiens]	0.0004
91	13031937	transcription factor i [Homo sapiens]	0.0004
		gi]14574479 gb AAF60869.2  (AC024856) Hypothetical	
92	14574479	protein Y71G10AR.2 [Caenorhabditis elegans]	0.15
	11371175	gi 14970562 emb CAC44371,1  (AJ292465) WDR9	0.15
93	14970562	protein, form A [Homo sapiens]	2E-16
		gi 12855751 dbj BAB30445.1  (AK016814) putative	22.10
98	12855751	[Mus musculus]	3.1
		gi 11498284 ref NP 069510.1  adenylate kinase (adk)	
		[Archaeoglobus fulgidus] sp Q29581 KAD ARCFU	
		ADENYLATE KINASE (ATP-AMP	
		TRANSPHOSPHORYLASE) pir D69334 adenylate	
		kinase (EC 2.7.4.3) - Archaeoglobus fulgidus	
		gb AAB90565.1  (AE001058) adenylate kinase (adk)	
102	11498284	[Archaeoglobus fulgidus]	4.00E-13
		gi 9910248 ref[NP 064579.1  GL004 protein [Homo	
		sapiens] gb AAF86949.1 AF226049 1 (AF226049)	
104	9910248	GL004 [Homo sapiens]	1.00E-85
		gi 7302191 gb AAF57287.1  (AE003784) CG7856 gene	
105	7302191	product [Drosophila melanogaster]	1.00E+00
		gi 12005513 gb AAG44486.1 AF246239_1 (AF246239)	
	1	AD030 [Homo sapiens] gb AAG44658.1 AF258660_1	
106	12005513	(AF258660) AD033 [Homo sapiens]	2.00E-73
		gi 13874435 dbj BAB46923.1  (AB000782) cerebral	
108	13874435	protein-10 [Homo sapiens]	4.00E-04
		gi 14783375 ref XP_045533.1  protein predicted by	
109	14783375	clone 23627 [Homo sapiens]	1.60E-01
		gi 7229532 gb AAF42865.1 AF225924_1 (AF225924)	
110	7229532	staufen [Drosophila virilis]	3.30E+00
110	14555001	gi 14775931 ref XP_049935.1  hypothetical protein	
112	14775931	FLJ14950 [Homo sapiens]	3.00E-77
		-1101407581-114-4-012580-114-0051-622-5	
114	10140750	gi 10140758 gb AAG13589.1 AC051633_5 (AC051633) putative ubiquitin protein [Oryza sativa]	2 10E±00
114	10140/38	[(ACO31033) putative ubiquitin protein [Oryza sativa]	2.10E+00

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 421058 pir  S34255 hypothetical protein 1a -	
		Escherichia coli emb CAA51036.1  (X72295) ORF1a	
115	421058	[Escherichia coli]	6.50E+00
		gi 14775931 ref XP_049935.1  hypothetical protein	
116	14775931	FLJ14950 [Homo sapiens]	2.00E-74
		gi 14781316 ref XP_032533.1  hypothetical protein	
117	14781316	FLJ12787 [Homo sapiens]	e-101
		gi 13161938 cmb CAC32991.1  (AJ309861) putative	
118	13161938	protein kinase WNK4 [Homo sapiens]	1.00E-30
		gi 7662028 ref[NP_055557.1  KIAA0255 gene product	
		[Homo sapiens] ref[XP_009540.1] KIAA0255 gene	
		product [Homo sapiens] sp Q92544 T9S4_HUMAN	
		TRANSMEMBRANE 9 SUPERFAMILY PROTEIN	·
		MEMBER 4 dbj BAA13385.1  (D87444) Similar to	
		S.cerevisiae EMP70 protein precursor (S25110) [Homo	
		sapiens] emb CAB75607.2  (AL049539) dJ836N17.2	
119	7662028	(KIAA0255 protein) [Homo sapiens]	7.00E-14
		gi 10179324 dbj BAB13674.1  (AB041351) type IV	
120	10179324	collagen alpha 6 chain [Mus musculus]	7.9
		gi 7160119 emb CAB76306.1  (AL158057) putative	
		acetyltransferase (fragment). [Streptomyces coelicolor	
122	7160119	A3(2)]	0.1
		gi 14732244 ref XP_039960.1  PC3-96 protein [Homo	
		sapiens] ref[XP 039961.1] PC3-96 protein [Homo	
		sapiens) pir T46276 hypothetical protein	
		DKFZp564M1178.1 - human emb CAB70781.1	
		(AL137515) hypothetical protein [Homo sapiens]	
		gb AAG35611.1 AF202092_1 (AF202092) PC3-96	
123	14732244	[Homo sapiens]	9E-81
		gi 7020475 dbj BAA91144.1  (AK000407) unnamed	
124	7020475	protein product [Homo sapiens]	2.6
		gi 14775931 ref XP_049935.1  hypothetical protein	
125	14775931	FLJ14950 [Homo sapiens]	2E-72
	l	gi 226135 prf  1411303A GABA receptor alpha2 [Bos	
127	226135	taurus]	0.046
		gi 4995818 emb CAB44313.1  (AJ131899) proline rich	
136	4995818	synapse associated protein 1 [Rattus norvegicus]	0.00003

	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS			
NO	N	DESCRIPTION	P VALUE	
	1			
	l	gi 13357869 ref[NP 078143.1  unique hypothetical		
		[Ureaplasma urealyticum] pir D82907 hypothetical		
	1	protein UU309 [imported] - Ureaplasma urealyticum		
	Ì	gb AAF30718.1 AE002128_6 (AE002128) unique		
138	13357869	hypothetical [Urcaplasma urealyticum]	2.7	
		gi 1076802 pir  S49915 extensin-like protein - maize		
		emb CAA84230.1  (Z34465) extensin-like protein [Zea		
139	1076802	mays] prf[2111476A extensin-like domain [Zea mays]	1.9	
		gi 11466208 ref NP 066531.1  NADH dehydrogenase		
	1	subunit 4 [Naegleria gruberi]		
	1	gb AAG17809.1 AF288092 34 (AF288092) NADH		
140	11466208	dehydrogenase subunit 4 [Naegleria gruberi]	6.6	
	11100200	gi 7490290 pir T38644 conserved hypothetical protein	0.0	
		SPAC323.07c - fission yeast (Schizosaccharomyces		
		pombe) emb CAB53410.1  (AL109988) conserved		
*	1	hypothetical protein; UPF0013 [Schizosaccharomyces		
145	7490290		0.78	
		gi 5813770 gb AAD52006.1 AF017304 1 (AF017304)		
147	5813770	FMVIB [Morone saxatilis]	6,8	
		gil2497311 sp P55803 MOG BOVIN MYELIN-		
	1	OLIGODENDROCYTE GLYCOPROTEIN		
		PRECURSOR pir  A47712 myelin/oligodendrocyte		
149	2497311	glycoprotein precursor - bovine	0.33	
		gi 609342 gb AAA58698.1  (U04946) nucleophosmin-		
	1	anaplastic lymphoma kinase fusion protein [Homo		
152	609342	sapiens]	2.7	
		gi 7481152 pir  T36795 probable penicillin acylase -		
	1	Streptomyces coelicolor emb CAB46792.1  (AL096811)		
		putative penicillin acylase [Streptomyces coelicolor		
153	7481152	A3(2)]	8	

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
154	12229694	gi 12229694 sp 093830 BET2_CANAL TYPE II PROTEINS GERANYLGERANYLTRANSFERASE BETA SUBUNIT (TYPE II PROTEIN GERANYL- GERANYLTRANSFERASE BETA SUBUNIT) (GGTASE-II-BETA) (PGGT) (YPTI/SEC4 PROTEINS GERANYLGERANYLTRANSFERASE BETA SUBUNIT) dbj BAA35193.I  (AB021171) Beta subunit of geranylgeranyl transferase type2 [Candida albicans]	4.7
156	0620053	gi 9630058 rcf NP_046276.1  unknown [Orgyia pseudotsugata single capsid nuclear polyhedrosis virus] sp 010359 Y120_NPVOP HYPOTHETICAL 9.3 KD PROTEIN (ORF120) pir T10389 hypothetical protein 120 - Orgyia pseudotsugata nuclear polyhedrosis virus gb AAC59119.1  (U75930) unknown [Orgyia	
156	9630058	pseudotsugata single capsid nuclear polyhedrosis virus] gil9965966 gb AAG10219.1 AF294433 1 (AF294433)	8.5
157	9965966	coat protein [Alfalfa mosaic virus]	7
158	7296047	gi 7296047 gb AAF51343,1  (AE003585) CG17711 gene product [Drosophila melanogaster]	4.6
161	13507856	gil 13507856[ref]NP_109805.1] ribosomal protein L20 [Mycoplasma pneumoniae] splP78023]RL20_MYCPN 50S RIBOSOMAL PROTEIN L20 pir[873363 ribosomal protein L20 - Mycoplasma pneumoniae (strain ATCC 29342) gb]AAG34734.1]AE000004_3 (AE000004) ribosomal protein L20 [Mycoplasma pneumoniae]	0.86
		gi 8393641 ref NP_058889.1  kynurenine aminotransferase II [Rattus norvegicus] emb CAA90507.1  (Z50144) kynurenine/alpha-	
162	8393641	aminoadipate aminotransferase [Rattus norvegicus]	4.2
169	1515448	gi 1515448 gb AAC45559.1  (U63096) Description: pBF4 gene involved in Bacteroides spp. conjugal transfer [Bacteroides fragilis]	2.6
175	422832	gi 422832 pir  B46629 mucin 6, gastric (3-repeat clone) - human (fragment) gb AAB61945.1  (L07518) mucin [Homo sapiens]	3.2

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS		
NU	N	DESCRIPTION	P VALUE
	1	gi 7504075 pir  T22584 hypothetical protein F53F4.6 -	
100	2504025	Caenorhabditis elegans emb CAB01215.1  (Z77663)	
176	7504075	F53F4.6 [Cacnorhabditis elegans]	6.6
		-1122955291 (BVD 09021 ( 11 DVCENT DATA	
		gi 13385538 ref NP_080316.1  RIKEN cDNA 2810036K01 gene [Mus musculus] dbi BAB28520.1	
178	12205520	(AK012865) putative [Mus musculus]	277.40
170	13363336	gi 902377 gb AAA82981.1  (U18059) polyprotein	2E-48
180	902377	[pestivirus type 1]	0.50
160	902377	ipestivirus type 1j	0.53
		gi 13421499 gb AAK22335.1  (AE005708) conserved	
181	13/21/00	hypothetical protein [Caulobacter crescentus]	6.2
101	13421499	hypothetical protein [Cautobacter crescentus]	0.2
		2100005741 WID 045070 H.C. NETT 1	
		gi 10803574 ref NP_045972.1  GvpN [Halobacterium sp. NRC-11 sp O9HI16 GVN1 HALN1 GVPN	
		PROTEIN 1 pir A47053 gas-vesicle operon protein	
		gvpN [imported] - Halobacterium salinarum pir  T08244 gas-vesicle operon protein gvpN - Halobacterium sp.	
		(strain NRC-1) plasmid pNRC100 gblAAC82811.1	
		(AF016485) GypN [Halobacterium sp. NRC-1]	
		gb AAD15044.1  (L03361) gas vesicle protein [Plasmid	
		pNRC100] gb AAG20728.1  (AE005142) GvpN	
187	10803574	protein, cluster A; GvpN1 [Halobacterium sp. NRC-1]	7.40E-01
10,	10003377	gi/7662532 refINP 054838.1  PRO0195 protein [Homo	7.4015-01
		sapiens] gb AAF24025.1 AF090901 1 (AF090901)	
192	7662532	PRO0195 [Homo sapiens]	2.60E-01
		gi 5360226 dbi BAA36472.1  (AB015177) F0-ATPase	2.002-01
194	5360226	subunit 6 [Beta vulgaris]	2.3
		gi 3024921 sp Q58312 Y902 METJA	
		HYPOTHETICAL PROTEIN MJ0902 pir  F64412	
		hypothetical protein MJ0902 - Methanococcus	
		jannaschii gb AAB98907,1  (U67533) M. jannaschii	
		predicted coding region MJ0902 [Methanococcus	
196	3024921	jannaschii]	6.5
		gi 7522108 pir T29097 pro-pol-dUTPase polyprotein -	
		murine endogenous retrovirus ERV-L (fragment)	
		cmb CAA73251.1  (Y12713) protease; reverse	
		transcriptase; RNaseH; integrase; dUTPase; Pro-Pol-	
198	7522108	dUTPase polyprotein [Mus musculus]	1.7

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SEQ ID   ACCESS   DESCRIPTION   P N	ALUE
	ALUE
ail0955592lomblCAC00652 1L(A1202510) N1	
ail0055502lomblCAC00652 11 (A1202510) N1	
199 9955592 acid racemase [Amycolatopsis orientalis subsp. lurida]	4.5
gi 4097932 gb AAD10321.1  (U72240) unknown	
203 4097932 [Choristoneura fumiferana nucleopolyhedrovirus]	6.5
gi 11252252 pir  D82079 pyruvate dehydrogenase, E1	
component VC2414 [imported] - Vibrio cholerae (group	
O1 strain N16961) gb AAF95557.1  (AE004311)	
pyruvate dehydrogenase, E1 component [Vibrio	
205   11252252   cholcrac]	3.5
gi 12963481 ref NP_061927.1  hypothetical protein	
MGC5560; hypothetical protein [Homo sapiens]	
dbj BAB14342.1  (AK022978) unnamed protein product	
	E-73
gi 14737646 ref XP_038048.1  similar to APICAL	
ENDOSOMAL GLYCOPROTEIN PRECURSOR (R.	
207   14737646   norvegicus) [Homo sapiens]	4
gi 14742770 ref XP_039393.1  KIAA1550 protein	0.4
209   14742770 [Homo sapiens]	8.4
gi 12643499 sp P89202 RRPO_SHMV RNA-	
DIRECTED RNA POLYMERASE (186 KDA PROTEIN) ICONTAINS:	
METHYLTRANSFERASE/RNA HELICASE	
213 12643499 (MT/HEL) (128 KDA PROTEIN)]	6.5
	0.3
gi 9629141 ref NP_044299.1  putative transport protein	
[Soybean chlorotic mottle virus]	
sp P15631 VMP_SOCMV MOVEMENT PROTEIN (CELL-TO-CELL TRANSPORT PROTEIN) (ORF	
IA) pirlJS0379 hypothetical 35.5K protein - soybean	
chlorotic mottle virus emb CAA33833.1  (X15828)	
putative transport protein [Soybean chlorotic mottle	
223 9629141 virus]	4.3
gi 7494266 pir  T18485 hypothetical protein C0840w -	
malaria parasite (Plasmodium falciparum)	
emb CAB11124,1  (Z98551) putative P-type ATPase	
226 7494266 [Plasmodium falciparum]	2.3
gi 14042283 dbi BAB55184.1  (AK027535) unnamed	
01 1-31	E-30
gi 14744326 ref XP 044812.1  NIMA (never in mitosis	
229 14744326 gene a)-related kinase 6 [Homo sapiens] 4	E-13
gi 98014 pir  S11148 amiA protein - Streptococcus	
232 98014 pneumoniae	10

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 10178317 emb CAC08403.1  (AL121928)	
233	10178317	bA18I14.5.1 (novel protein isoform 1) [Homo sapiens]	7
1	•	gi 630444 pir  S43955 probable NADH dehydrogenase	
		(ubiquinone) (EC 1.6.5.3) chain 3, kinetoplast -	
		Trypanosoma brucei mitochondrion (fragment)	
234	630444	gb AAA20887.1  (L26251) CR5 [Trypanosoma brucci]	6.2
		gi 7505826 pir  T25816 hypothetical protein K12D9.2 -	
239	7505826	Caenorhabditis elegans	1.4
		gi 7434912 pir  H71934 phosphatidylglycerophosphate	
l		synthase - Helicobacter pylori (strain J99)	
1	ŀ	gb AAD05990.1  (AE001475)	
		PHOSPHATIDYLGLYCEROPHOSPHATE	
240	7434912	SYNTHASE [Helicobacter pylori J99]	5.8
	,	gi 6136155 sp O70546 UTX_MOUSE	
	ļ	UBIQUITOUSLY TRANSCRIBED X	
	l	CHROMOSOME TETRATRICOPEPTIDE REPEAT	
		PROTEIN (UBIQUITOUSLY TRANSCRIBED TPR	
		PROTEIN ON THE X CHROMOSOME)	
243	6136155	emb CAA05692.1  (AJ002730) UTX [Mus musculus]	8.90E+00
1			
		gi 2143962 pir  I59422 rsec8 - rat (fragment)	
246	2143962	gb AAC52265.1  (U32498) rsec8 [Rattus norvegicus]	1E-93
	'		
		gi 6136155 sp O70546 UTX_MOUSE	
1	1	UBIQUITOUSLY TRANSCRIBED X	
		CHROMOSOME TETRATRICOPEPTIDE REPEAT	
		PROTEIN (UBIQUITOUSLY TRANSCRIBED TPR	
	l	PROTEIN ON THE X CHROMOSOME)	
250	6136155	emb CAA05692.1  (AJ002730) UTX [Mus musculus]	7.1

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteine)
SEO ID	ACCESS	Table 55 Treatest Treigness (Blaster Vs. Troir-Rediffication	T Totolis)
NO	N	DESCRIPTION	P VALUE
İ			
		gi 12737603 ref XP 006839,2  activin A receptor type II	
		like 1 [Homo sapiens] ref[XP 050707.1] activin A	
	ì	receptor type II-like 1 [Homo sapiens]	1
		sp P37023 KIR3_HUMAN SERINE/THREONINE-	
		PROTEIN KINASE RECEPTOR R3 PRECURSOR	
		(SKR3) (ACTIVIN RECEPTOR-LIKE KINASE 1)	
	ļ	(ALK-1) (TGF-B SUPERFAMILY RECEPTOR TYPE	
	i	I) (TSR-I) pir  A49431 activin/TGF-beta-like type I	
		receptor - human gb AAA16160.1  (L17075) TGF-b	
		superfamily receptor type I [Homo sapiens]	
		gb AAB61900.1  (U77713) activin receptor like kinase 1	
255	12737603	[Homo sapiens]	7.50E+00
		gi 2133974 pir  S68425 SoxP1 protein - rainbow trout	
		dbj[BAA11868.1] (D83256) SoxP1 [Oncorhynchus	
265	2133974		0.18
266	9838427	gi 9838427 ref NP_064041.1  orf214 [Bcta vulgaris]	2.2
268	5721727	gi 5731737 dbj BAA83337.1  (AB021878) similar to	2.2
208	5731737	yeast sodium/proton exchanger [Oryza sativa]	3,3
		gi 13642312 ref XP_002895.2  parathyroid hormone	
	i	receptor 1 [Homo sapiens] ref[XP_033742.1]	
		parathyroid hormone receptor 1 [Homo sapiens] ref[XP 033743.1  parathyroid hormone receptor 1	
271	12642212	[Homo sapiens]	9
	13042312	gi 11545138 cmb CAC08390.2  (AL121886)	
		dJ1028D15,1 (continued from dJ138B7,1 in	
272	11545138	Em:Z98752) [Homo sapiens]	0,0000008
	11313130	Enn. 25 0 ( 52) [Nono superis]	0,000000
		gi 6323677 ref[NP 013748,1  Ymr034cp	
		[Saccharomyces cerevisiae] splO05131[YMS4 YEAST	
		HYPOTHETICAL 48.4 KD PROTEIN IN ARP9-	
		IMP2 INTERGENIC REGION pir  S53951 probable	
		membrane protein YMR034c - yeast (Saccharomyces	
		cerevisiae) emb CAA89150.1  (Z49213) unknown	
275	6323677	[Saccharomyces cerevisiae]	6.6
		gi 7297043 gb AAF52312.1  (AE003612) CG13992	
277	7297043	gene product [Drosophila melanogaster]	8.1
		gi 1616595 emb CAA41710.1  (X58907) steroid 21-	
282	1616595	monooxygenase [Homo sapiens]	6.8
		gi 14089610 emb CAC13370.1  (AL445563) unknown;	
283	14089610	predicted coding region [Mycoplasma pulmonis]	2.1

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 139809 sp P27571 XIST_MOUSE X INACTIVE	
		SPECIFIC TRANSCRIPT PROTEIN pir   S15433	
1		hypothetical protein - mouse emb CAA41978.1	l .
		(X59289) ORF [Mus musculus] prf  1711440A xist	
284	139809	gene [Mus musculus]	0.18
		gi 11355576 pir  T44331 hypothetical protein wblD	
		[imported] - Vibrio cholerae dbj BAA33635.1	
		(AB012957) probable NADH dehydrogenase [Vibrio	
285	11355576	cholerae]	0.23
		gi 14389443 ref[NP_116776.1  NADH dehydrogenase	
		subunit 4 [Rana nigromaculata] dbj BAB58996.1	
		(AB043889) NADH dehydrogenase subunit 4 [Rana	
297	14389443	nigromaculata]	9.2
		gi 4493987 emb CAB39046.1  (AL034559) hypothetical	
300	4493987	protein, PFC1030w [Plasmodium falciparum]	2.1
		gi 7494200 pir  T18434 hypothetical protein C0375c -	
		malaria parasite (Plasmodium falciparum)	
		emb CAB11111.1  (Z98547) predicted using hexExon;	
		MAL3P3.10 (PFC0375c), Homologue of C.elegans	
		T08A11.2 protein, len: 1387 aa; Similarity to C.clegans	
		T08A11.2 protein. C.clegans T08A11.2 protein	
200		(WP:T08A11.2) BLAST Score: 2901, sum P(2) = 0.0;	
306	7494200	66% identity in 839 aa >	1.3
200	10700064	gi 10728064 gb AAF50455,2  (AE003556) CG7060	
308	10728064	gene product [Drosophila melanogaster]	4.7
		gi 9631682 ref[NP_048461.1  a113L [Paramecium	
1		bursaria Chlorella virus 1] pir  T17603 hypothetical	
1		protein al 13L - Chlorella virus PBCV-1	
217	0621682	gb AAC96481.1  (U42580) a113L [Paramecium	
317	9631682	bursaria Chlorella virus 1]	7.7
}		gi 4733986 gb AAD28666.1 AC007209_2 (AC007209)	
318	4733986	hypothetical protein [Arabidopsis thaliana]	5.30E+00
316	+133960	nypomenear protein [Arabidopsis tranana]	3.30E+00
		-: 14616401- 1005004IDDG1 BAREEDBUGH BORDER	
		gi 461649 sp Q05004 BB61_RABIT BRUSH BORDER 61.9 KD PROTEIN PRECURSOR pir B45665 adult-	
		specific 61.9K brush border protein precursor - rabbit	
		emb CAA78302.1  (Z12840) protein of unknown	
319	461649	function [Oryctolagus cuniculus]	3,00E-16
317	701049	ranction (Oryctolagus cuinculus)	3,00E-10

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID NO	ACCESS N	DESCRIPTION	DIVITE	
110	IN .	DESCRIPTION	P VALUE	
l	Į			
	1	gi 11466552 reflNP 044801.1  NADH dehydrogenase.		
	i	subunit 4 [Reclinomonas americana] pir S78183 NADH		
		dehydrogenase (ubiquinone) (EC 1.6.5.3) chain 4 -		
i	1	Reclinomonas americana (ATCC 50394) mitochondrion		
		gb AAD11916.1  (AF007261) NADH dehydrogenase,		
322	11466552	subunit 4 [Reclinomonas americana]	9.1	
323	10.00000	gi 13603897 gb AAK31986.1 AF295594_1 (AF295594)		
323	13603897	SKTx1 precursor [Buthus martensii]	9.3	
	Ì	"ILLANGUEGI" (DATA OCCORDO ALLA ALL'ALLA		
		gi 11497767 rcf NP_068989.1  hypothetical protein [Archaeoglobus fulgidus] sp O30087 Y150 ARCFU		
i		HYPOTHETICAL PROTEIN AF0150 pir  F69268		
		hypothetical protein AF0150 - Archaeoglobus fulgidus		
		gb AAB91087.1  (AE001096) A. fulgidus predicted		
325	11497767	coding region AF0150 [Archaeoglobus fulgidus]	7.1	
	l	gi 7506939 pir  T24356 hypothetical protein T02D1.3 -		
		Caenorhabditis elegans emb CAB05908.1  (Z83319)		
332	7506939	T02D1.3 [Caenorhabditis elegans]	3.9	
1				
l		gi 7510675 pir  T29689 hypothetical protein ZC266.2 -		
336	7510675	Caenorhabditis elegans gb AAA96101.1  (U53140) ZC266.2 gene product [Caenorhabditis elegans]	2.4	
330	7310073	gi 12697923 dbi BAB21780.1  (AB051476) KIAA1689	2.4	
337	12697923	protein [Homo sapiens]	3E-21	
		E [ parkyone]		
		gi 630465 pir  S47042 protein kinase (EC 2.7.1.37) cdc2-		
338	630465	related 1 - malaria parasite (Plasmodium falciparum)	5.7	
		gi 7488318 pir T01191 RNA-directed DNA polymerase		
		homolog F21E10.5 - Arabidopsis thaliana		
1		gb AAC13599.1  (AF058914) similar to reverse		
		transcriptase (Pfam: transcript_fact.hmm, score: 72.31)		
340	7488318		5	
1		gi 5804818 cmb CAB52872.1  (AL021712) putative		
341	5804818	protein [Arabidopsis thaliana] emb CAB79171.1  (AL161556) putative protein [Arabidopsis thaliana]	5	
341	2004018	gil14579376 gb AAK69253.1 AF336309 48		
343	14579376	(AF336309) SpyB [Yersinia enterocolitica]	8.2	
2.2	5.5570	( seeses, spjb [ a dismin distribution]	0.2	

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	
NO	IN	DESCRIPTION	P VALUE
		gi 11284162 pir  H81077 hypothetical protein NMB1490	
		[imported] - Neisseria meningitidis (group B strain	
ì ·		MD58) gb AAF41846.1  (AE002498) hypothetical	!
344	11284162	protein [Neisseria meningitidis MC58]	3
		gi 11361512 pir H82781 hypothetical protein XF0626	
		[imported] - Xylella fastidiosa (strain 9a5c)	
}		gb AAF83436.1 AE003908_4 (AE003908) hypothetical	
347	11361512	protein [Xylella fastidiosa 9a5c]	7.6
1		gi 7497054 pir  T19739 hypothetical protein C35A5.4 -	
		Caenorhabditis elegans emb CAA94905.1  (Z71185)	
		contains similarity to Pfam domain: PF00635 (MSP (Major sperm protein) domain), Score=66.4, E-value=2e	
348	7497054	[Major sperm protein) domain), Score=66.4, E-value=26 16, N=1 [Caenorhabditis elegans]	0.064
340	1491034	gil6531601lgblAAF15500.1l (AF167672)	0.004
		immunoglobulin heavy chain variable region [Homo	
349	6531601	sapiens	1.6
		gi 12858471 dbj BAB31327.1  (AK018655) putative	
352	12858471	[Mus musculus]	3.9
		gi 2128157 pir  C64319 hypothetical protein MJ0154 -	
353	2128157	Methanococcus jannaschii	0.98
		gi 14195575 sp P58114 YV78_CAUCR	
		HYPOTHETICAL PROTEIN CC3178	
355	14105575	gb AAK25140.1  (AE005981) pirin-related protein [Caulobacter crescentus]	4 2072   00
. 333	14193373	[Caulonacter crescentus]	4.20E+00
		- 1174 C 842 5  - 1177 200 1 5   11	
		gi 7468435 pir B72015 metalloproteinase, insulinase family CP0903 [imported] - Chlamydophila pneumoniae	
		(strains CWL029 and AR39) gb AAD19093.1	
		(AE001675) Insulinase family/Protease III	
		[Chlamydophila pneumoniae CWL029]	'
		gb AAF38689.1  (AE002249) metalloprotease,	
l		insulinase family [Chlamydophila pneumoniae AR39]	
		dbj BAA99165.1  (AP002548) insulinase	
359	7468435	family/protease III [Chlamydophila pneumoniae J138]	4.1
		gi 461649 sp Q05004 BB61_RABIT BRUSH BORDER	
1 .		61.9 KD PROTEIN PRECURSOR pir B45665 adult	
'		specific 61.9K brush border protein precursor - rabbit emb CAA78302.1  (Z12840) protein of unknown	
360	461649	function [Oryctolagus cuniculus]	8E-45
500	401049	minorion [OTYGORIGUS CHIRCHIUS]	0L-43

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
SEQ ID	ACCESS			
NO	N	DESCRIPTION	P VALUE	
		gi 12859685 dbj BAB31736.1  (AK019464) putative		
363	12859685	[Mus musculus]	6.1	
	l			
		gi 11055982 ref NP_067633.1  PR domain containing		
		13; PR-domain containing protein 13 [Homo sapiens]		
		ref[XP_011460.1] PR domain containing 13 [Homo		
	ŀ	sapiens] gb AAG13448.1  (AY004253) PR-domain		
365	11055982	containing protein 13 [Homo sapiens]	3.4	
		gi 3915305 sp Q96597 VP5_AHSV9 OUTER CAPSID		
		PROTEIN VP5 gb AAB17570.1  (U74489) VP5		
368	3915305	[African horse sickness virus]	0.63	
i I	l	gi 11466986 ref NP_041893.1  Orf635 [Euglena	,	
		gracilis] sp P31916 YCX3_EUGGR HYPOTHETICAL		
		78.0 KD PROTEIN IN PSBC INTRON 2 (ORF635)		
370	11466986	emb CAA50080.1  (X70810) Orf635 [Euglena gracilis]	1.7	
		gi 14766918 ref XP_034599.1  acid sphingomyelinase-		
373	14766918	like phosphodiesterase [Homo sapiens]	2E-99	
		gi 5732610 gb AAD49190.1 AF115423_2 (AF115423)		
375	5732610	core protein [Hepatitis B virus]	3.4	
		gi 2707661 gb AAB94699.1  (AF029934) IgM heavy		
377	2707661	chain VDJ region [Oryctolagus cuniculus]	4.7	
		gi 7516819 pir  C72580 hypothetical protein APE1923 -		
		Aeropyrum pernix (strain K1) dbj BAA80928.1		
ĺ		(AP000062) 105aa long hypothetical protein		
379	7516819	[Aeropyrum pernix]	6.80E+00	
	1	gi 3859670 emb CAA22008.1  (AL033502) hypothetical		
380	3859670	membrane protein [Candida albicans]	9.90E+00	
		gi 14784562 ref XP_040415.1  mutL (E. coli) homolog 3		
	<u>-</u>	[Homo sapiens] gb AAF23905.1  (AF195658) DNA		
382	14784562	mismatch repair protein [Homo sapiens]	2E-25	
		gi 14194519 sp Q9K920 CMGB_BACHD COMG		
		OPERON PROTEIN 2 HOMOLOG dbj BAB06550,1		
l		(AP001516) DNA transport machinery [Bacillus		
383	14194519	halodurans]	2.7	
		gi 7509624 pir  T26676 hypothetical protein Y38F1A.1 -		
		Caenorhabditis elegans emb CAA21628.1  (AL032639)		
		predicted using Genefinder~cDNA EST		
	2500504	EMBL:AW057282 comes from this gene		
386	7509624	[Caenorhabditis elegans]	9.4	

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 7508443 pir  T25281 hypothetical protein T25E12,11	
	1	Caenorhabditis elegans emb CAB04829.1  (Z82052)	
		contains similarity to Pfam domain: PF01827 (Domain	
388	7500440	of unknown function), Score=130.6, E-value=4.6e-36,	
300	7508443	N=1 [Caenorhabditis elegans] gi 7295832 gb AAF51133,1  (AE003580) CG3347 gene	3.4
389	7295832		0.022
389	1293832	product [Drosophila melanogaster] gi 7510359 pir  T27283 hypothetical protein Y64G10A.f	0.032
390	7510359		0.0
390	7510359		8.2
	1	gi 6679381 ref[NP_032903.1  plasminogen [Mus	
		musculus] sp[P20918]PLMN_MOUSE	
	l	PLASMINOGEN PRECURSOR [CONTAINS: ANGIOSTATIN] pir  PLMS plasmin (EC 3.4.21.7)	
		precursor - mouse gblAAA50168.11 (J04766)	
395	6679381	plasminogen [Mus musculus]	2.2
393	0079381	prasmmogen (ivius musculus)	2.2
		gi 8479518 sp Q89853 VGP_EBORS STRUCTURAL	
		GLYCOPROTEIN PRECURSOR (VIRION SPIKE	
		GLYCOPROTEIN) [CONTAINS: GP1; GP2] gb AAC54889.1  (U23416) virion spike glycoprotein	
		[Ebola virus] gb AAC54891.1  (U23417) virion spike	
396	0470510	glycoprotein [Ebola virus]	3.1
390	04/9310	gi 14133247 dbi BAA86564.2  (AB033076) KIAA1250	3.1
403	14122247	protein [Homo sapiens]	6E-19
403	14133247	gi 3293234 gb AAC25913.1  (U96413) T-DNA	0E-19
412	2202224	oncoprotein [Agrobacterium tumefaciens]	9.1
412	3293234	gi 7023033 dbi BAA91809.1  (AK001649) unnamed	9.1
415	7023033	protein product [Homo sapiens]	9E-85
713	7023033	protein product [Fromo sapiens]	912-03
		gi 12514130 gb AAG55434,1 AE005285 1 (AE005285)	
		putative oxidoreductase [Escherichia coli O157:H7	
		EDL933] dbi BAB34455,1  (AP002554) putative	
416	12514130	oxidoreductase [Escherichia coli O157:H7]	9.40E-01
710	12314130	gi 4514359 dbi BAA75394.1  (AB013377) ComGB	7,700-01
418	4514359	[Bacillus halodurans]	2.6
		gi 7510324 pir  T27250 hypothetical protein Y5F2A.3 -	
		Caenorhabditis elegans emb CAA21648.1  (AL032641)	
419	7510324	Y5F2A.3 [Cacnorhabditis elegans]	1.6

	Γ	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteine)
SEO ID	ACCESS	Table 55 Homest Heighbor (BlastA Vs. Hon-Redifficant	1 Towns)
NO	N	DESCRIPTION	P VALUE
	1	gi 6981358 ref NP 037137,1  phosphoinositide 3-kinase	
		p85 (other splicing variants: p55 and p50) [Rattus	
	1	norvegicus] sp Q63787 P85A RAT	
	1	PHOSPHATIDYLINOSITOL 3-KINASE	
		REGULATORY ALPHA SUBUNIT (PI3-KINASE	
	1	P85-ALPHA SUBUNIT) (PTDINS-3-KINASE P85-	
		ALPHA) (PI3K) dbj BAA18932.1  (D64045)	
		phosphatidylinositol 3-kinase p85 alpha subunit [Rattus	
421	6981358	norvegicus]	2.4
		gi 6576738 dbj BAA88337.1  (AB005891) ORF2	
423	6576738	[Platemys spixii]	3.4
		gi 6739553 gb AAF27299.1  (AF146429) DeltaC [Danio	
424	6739553	rerio]	3.9
		gi 13816283 gb AAK43020.1  (AE006883) Sulfate	
425	13816283	adenylyltransferase (sat) [Sulfolobus solfataricus]	3.6
		  gi 11034774 gb AAG27071,1 AF105225 7 (AF030414)	
428	11024774	NifQ [Gluconacetobacter diazotrophicus]	3.6
420	11034774	gi 13364676 dbj BAB38622.1  (AP002568) hypothetical	3.0
433	12264676	protein [Escherichia coli O157:H7]	5,8
433	13304070	gi 12845784 dbi BAB26898.1  (AK010380) putative	3,6
434	12845784		0.075
7,57	12043704	[ivius muscalus]	0.073
		  gi 12513917 gb AAG55266.1 AE005270_2 (AE005270)	
		Z1121 gene product [Escherichia coli O157:H7	
		EDL933] gb AAG55675.1 AE005306 4 (AE005306)	
		Z1560 gene product [Escherichia coli O157:H7	
		EDL933] dbj BAB34723.1  (AP002554) putative	
435	12513917	membrane protein [Escherichia coli O157:H7]	8.3
	12515717	gi/7661654/ref NP 056444.1  DKFZP566J153 protein	
		[Homo sapiens] emb[CAB43677.1] (AL050369)	
436	7661654	hypothetical protein [Homo sapiens]	0,42
	7001051	gi 1150678 emb CAA50971.1  (X72086) ORF20R;	0,12
437	1150678	B21R in citation [3] [Variola virus]	5.10E+00
		[7]	
	i	gi 8928456 sp O51039 Y006 BORBU	
		HYPOTHETICAL PROTEIN BB0006 pir F70100	
		conserved hypothetical integral membrane protein	
		BB0006 - Lyme disease spirochete gb[AAC66397.1]	
		(AE001115) conserved hypothetical integral membrane	
438	8928456	protein [Borrelia burgdorferi]	4.2
		gi 7297273 gb AAF52536.1  (AE003618) Myo28B1	
440	7297273	gene product [Drosophila melanogaster]	3.3

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	8-10-10-10-10-10-10-10-10-10-10-10-10-10-	11000000)
NO	N	DESCRIPTION	P VALUE
		gi 7495690 pir  T19090 hypothetical protein C08F11.3 -	
		Caenorhabditis elegans emb CAB05674.1  (Z83216)	
448	7495690	C08F11.3 [Caenorhabditis elegans]	4.9
		gi 10438579 dbj BAB15281.1  (AK025916) unnamed	
449	10438579		3.7
		gi 112200 pir  A32868 prolactin receptor - rat	
451	112200	(fragments)	8.6
1	ŀ	gi 114987 sp P17885 BIMA_EMENI BIMA PROTEIN	
		pir  A53256 nuclear protein bimA - Emericella nidulans	
		emb CAA41959.1  (X59269) bimA [Emericella	
452	114987	nidulans]	6.60E+00
		gi 4885039 gb AAD31932.1 U00058_6 (U00058)	
		contains similarity to DNAJ domains (Pfam: PF00226,	
457	4885039	Score=44.9, E=1.8e-09, N=1) [Caenorhabditis elegans]	4.3
		( iii )	
	1	gi 4505121 ref NP_003916.1  methyl-CpG binding	
		domain protein 4; 3,N(4)-ethenocytosine glycosylase;	
		G/T mismatch glycosylase; G/U mismatch glycosylase;	
		G/5-fluorouracil mismatch glycosylase with biphasic	
		kinetics [Homo sapiens] gb AAC68879.1  (AF072250)	
		methyl-CpG binding protein MBD4 [Homo sapiens]	
	1	gb AAD22195.1 AF114784_1 (AF114784) methyl-CpG binding endonuclease [Homo sapiens] gb AAD50374.1	
		(AF120999) methyl-CpG binding protein 4 [Homo	
459	4505121	sapiens]	2.1
1,57	1505121	gi 12644455 sp O60751 IG1R_MOUSE INSULIN-	2.1
		LIKE GROWTH FACTOR I RECEPTOR	
		PRECURSOR gb AAC12782.1  (AF056187) insulin-	
		like growth factor I receptor; IGF-I receptor [Mus	
463	12644455	musculus]	3.1
		gi 11465398 ref[NP 045211.1  unknown; N-acetyl-	
		glutamate-gamma-semialdehyde dehydrogenase	
		[Cyanidium caldarium] sp Q9TLQ8 HIS5 CYACA	
Ì	1	AMIDOTRANSFERASE HISH	
		gb AAF12883.1 AF022186_5 (AF022186) unknown; N-	
		acetyl-glutamate-gamma-semialdehyde dehydrogenase	
465	11465398	[Cyanidium caldarium]	
		gi 5761329 dbj BAA83473.1  (AB004819) cysteine	
466	5761329	endopeptidase [Oryza sativa]	5.70E+00

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
ĺ	İ		
		gi 5306243 gb AAD41976.1 AC006438_8 (AC006438)	
468	5306243	hypothetical protein [Arabidopsis thaliana]	4.2
		gi 7304025 gb AAF59067.1  (AE003836) CG8639 gene	
470	7304025	product [Drosophila melanogaster]	3.20E-01
}		gi 7303724 gb AAF58773.1  (AE003829) lola gene	
		product [alt 2] [Drosophila melanogaster]	
		gb AAF58774.1  (AE003829) lola gene product [alt 3]	
	l	[Drosophila melanogaster] gb AAF58775.1	
	l	(AE003829) lola gene product [alt 4] [Drosophila	
		melanogaster] gb AAF58776.1  (AE003829) lola gene	
473	7303724	product [alt 5] [Drosophila melanogaster]	5.8
		gi 1098549 gb AAA82594.1  (U25703) immunoglobulin	_
476	1098549	light chain F class [Ictalurus punctatus]	7
		gi 4262231 gb AAD14524.1  (AC006200) hypothetical	
477	4262231	protein [Arabidopsis thaliana]	3.8
		gi 5834894 ref NP_006964,1 ND5_10021 NADH	
		dehydrogenase subunit 5 [Caenorhabditis elegans]	
		sp[P24896]NU5M_CAEEL NADH-UBIQUINONE	
		OXIDOREDUCTASE CHAIN 5 pir    S26037 NADH	
		dehydrogenase (ubiquinone) (EC 1.6.5.3) chain 5 -	
		Caenorhabditis elegans mitochondrion	
.=0		emb CAA38162.1  (X54252) ND5 protein (AA 1 - 527)	
478	5834894	[Caenorhabditis elegans]	0.23
400	14776600	gi 14776608 ref XP_033588.1  BAI1-associated protein	
480	14776608	3 [Homo sapiens]	8.6
		gi 7448960 pir  D72417 conserved hypothetical protein -	
		Thermotoga maritima (strain MSB8) gb AAD35211.1 AE001697 11 (AE001697) conserved	
482	7448960	hypothetical protein [Thermotoga maritima]	0.1
402	7448900	nypotheticai protein [Thermotoga mariuma]	9.1
		**************************************	
		gi 7497960 pir  T15840 hypothetical protein C54G7.3 -	
		Caenorhabditis elegans gb AAA81392.1  (U40410)	
		coded for by C. elegans cDNA yk9e10.5; coded for by C. elegans cDNA yk9c10.3; multiple regions of	
		similarity to EGF-like repeats and cysteine-rich repeats	
484	7497960		5
704	1791900	gi 14720884 ref XP 032180.1  hypothetical protein	
487	14720884	DKFZp434A171 [Homo sapiens]	7.7
707	1-7720004	gi 3201900 gb AAC19365.1  (AF067420) SNC73	1.1
490	3201900	protein [Homo sapiens]	0.28
490	2201900	protein (riono sapiens)	0.28

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 12018149 gb AAG45421.1 AF309495 1 (AF309495)	
		gamete-specific hydroxyproline-rich glycoprotein a2	
493	12018149	[Chlamydomonas reinhardtii]	3.7
		gi 467634 emb CAA82856.1  (Z29969) HLY 4	
494	467634	[Entamoeba histolytica]	8.9
		gi 14726914 ref XP_037105.1  KIAA0622 protein;	
		Drosophila 'multiple asters' (Mast)-like homolog 1	
498	14726914	[Homo sapiens]	6.8
		gi 11692583 gb AAG39888.1 AF282303_1 (AF282303)	
499	11692583	odorant receptor M34 [Mus musculus]	9.5
		gi 7303064 gb AAF58132.1  (AE003810) CG12960	
502	7303064	gene product [Drosophila melanogaster]	4.9
		gi 13701254 dbj BAB42549.1  (AP003133)	
		ORFID:SA1289~hypothetical protein, similar to	
		bifunctional biotin ligase/biotin operon repressor	
		[Staphylococcus aureus subsp. aureus N315]	
		dbj BAB57618.1  (AP003362) hypothetical protein	
504	13701254	[Staphylococcus aureus subsp. aureus Mu50]	5.9
		gi 137073 sp P17086 URE1 PROMI UREASE ALPHA	
		SUBUNIT (UREA AMIDOHYDROLASE)	
		pir D43719 urease (EC 3.5.1.5) 62K chain - Proteus	
		mirabilis gb AAA25669.1  (M31834) urease subunit C	
505	137073	[Proteus mirabilis]	8.5
		gi 14548038 sp Q9D952 EVPL_MOUSE	
		ENVOPLAKIN (P210) (210 KDA CORNIFIED	
		ENVELOPE PRECURSOR) emb CAC38864.2	
512	14548038	(AJ309317) envoplakin [Mus musculus]	4.2
		gi 14031018 gb AAK50523.1  (AY029684) NADH	
516	14031018	dehydrogenase F [Stenotaphrum secundatum]	4.6
		gi 14600836 ref NP_147359.1  hypothetical protein	
		[Aeropyrum pernix] pir  D72647 hypothetical protein	
		APE0610 - Aeropyrum pernix (strain K1)	
		dbj BAA79580.1  (AP000060) 376aa long hypothetical	
518	14600836	protein [Aeropyrum pernix]	2.8

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 4930062 pdb 1MAA D Chain D, Mouse	
		Acetylcholinesterase Catalytic Domain, Glycosylated	
		Protein pdb 1MAA A Chain A, Mouse	
		Acetylcholinesterase Catalytic Domain, Glycosylated	
		Protein pdb 1MAA C Chain C, Mouse	
		Acetylcholinesterase Catalytic Domain, Glycosylated	
		Protein pdb 1MAA B Chain B, Mouse	
		Acetylcholinesterase Catalytic Domain, Glycosylated	
519	4930062	Protein	8.8
		gi 9757603 dbj BAB08147.1  (AB030877) maturase	
522	9757603	[Lilium mackliniae]	0.62
		gi 7445887 pir  T07052 probable potassium channel	
		protein SKT2 - potato emb[CAA70870.1] (Y09699) putative inward rectifying potassium channel [Solanum	
526	7445887	tuberosum	2.1
326	7443667	tuberosumj	2.1
		gi 2897832 dbi BAA24910,1  (AB000134) cytochrome	
528	2897832	oxidase subunit I [Prorocentrum micans]	9.8
320	2071032	Oxidase subuliit i [i forocentrum micans]	7.0
		gi 7446379 pir  T14039 protein kinase (EC 2.7.1.37),	
		myotonic dystrophy-associated - rat gb AAC02941.1	
		(AF021935) mytonic dystrophy kinase-related Cdc42-	
537	7446379	binding kinase [Rattus norvegicus]	8E-52
	7110577	gi 14779648 refIXP 027140.1  hypothetical protein	- 02 02
544	14779648	FLJ23239 [Homo sapiens]	6.3
		gi 7292503 gb AAF47906.1  (AE003481) CG15023	
546	7292503	gene product [Drosophila melanogaster]	6.3
		gi 4758758 ref[NP 004529.1  nucleosome assembly	
		protein 1-like 3 [Homo sapiens]	
		sp Q99457 NPL3_HUMAN NUCLEOSOME	
		ASSEMBLY PROTEIN 1-LIKE 3 dbj BAA08904,1	
548	4758758	(D50370) nucleosome assembly protein [Homo sapiens]	1.6
		gi 14530412 emb CAC42291.1  (Z69360) cDNA EST	
		EMBL:U52071 comes from this gene [Caenorhabditis	
554	14530412	elegans]	7
		gi 6680788 ref NP_031576.1  Bloom syndrome protein	
		homolog (human) [Mus musculus]	
		sp O88700 BLM_MOUSE BLOOM'S SYNDROME	
	5500,000	PROTEIN HOMOLOG emb CAB10933.1  (Z98263)	4.0
559	0680788	BLM protein [Mus musculus]	4.8

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#### Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins) SEO ID ACCESS NO N DESCRIPTION P VALUE gil3294495lgblAAC25822.11 (AF038608) contains similarity to Mus musculus tumor susceptibility protein 560 3294495 TSG101 (GB:U52945) [Caenorhabditis elegans] 0.78 gi|6319273|ref|NP_009356.1| pre-tRNA processing; Pta1p [Saccharomyces cerevisiae] sp|Q01329|PTA1 YEAST PTA1 PROTEIN pir S31299 pre-tRNA processing protein PTA1 - yeast (Saccharomyces cerevisiae) gb|AAA34919.1| (M95673) pta1 [Saccharomyces cerevisiae] gb[AAC04988,1] (U12980) Pta1p: Pre-tRNA processing involved protein 561 6319273 [Saccharomyces cerevisiae] 9.9 gi|5817732|gb|AAD52875.1|AF142703 1 (AF142703) 567 5817732 maturase-like protein [Ophrestia radicosa] 6.6 gil126296|splP08548|LIN1 NYCCO LINE-1 REVERSE TRANSCRIPTASE HOMOLOG prf||1207289B reverse transcriptase related protein 126296 [Nycticebus coucang] 569 0.00000002 gi|13816099|gb|AAK42875.1| (AE006870) Hypothetical 572 13816099 protein [Sulfolobus solfataricus] 2.1 gil12513361|gb|AAG54834.1|AE005228 1 (AE005228) putative glutaminase [Escherichia coli O157:H7 EDL933] dbj|BAB33961.1| (AP002552) putative 574 12513361 glutaminase [Escherichia coli O157:H71 3.3 gi|11350453|pir|B82965 hypothetical protein PA5456 [imported] - Pseudomonas acruginosa (strain PAO1) gblAAG08841.1lAE004958 11 (AE004958) 575 11350453 hypothetical protein [Pseudomonas aeruginosa] 9 gil1174945|sp|P43109|VEXB_SALTI_VI POLYSACCHARIDE EXPORT INNER-MEMBRANE PROTEIN VEXB pirlG36892 Vi polysaccharide capsule transporter VexB - Salmonella typhi dbj|BAA03197.1| (D14156) Wzm protein 1174945 [Salmonella typhi] 584 45 gj/4566616|gb|AAD23408.1|AF113531 1 (AF113531) follicle-stimulating hormone receptor precursor 586 4566616 [Coturnix coturnix] 2 gi|14746756|ref|XP 039102.1| similar to hypothetical 589 14746756 protein FLJ20378 (H. sapiens) [Homo sapiens] 0.007

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESS				
NO	N	DESCRIPTION	P VALUE		
	l	gi 7429823 pir  D69991 conserved hypothetical protein			
		yteU - Bacillus subtilis gb AAC00275.1  (AF008220)			
1		YteU [Bacillus subtilis] emb CAB14987.1  (Z99119)			
		similar to hypothetical proteins from B. subtilis			
590	7429823	[Bacillus subtilis]	0.49		
		gi 11877309 emb CAC19023.1  (AJ278707) ORF 4			
592	11877309	[Neisseria meningitidis phage 2120]	9.2		
		gi 14193715 gb AAK56102.1 AF332074_1 (AF332074)			
1		peroxisome proliferator-activated receptor binding			
594	14193715	protein [Mus musculus]	0.000002		
l		gi 14625344 gb AAK71419.1 U80842_9 (U80842)			
		Hypothetical protein ZC239.19 [Caenorhabditis			
595	14625344	elegans]	1.9		
		gi 1107614 emb CAA62696.1  (X91351) viral infectivity			
596	1107614	factor protein [Human immunodeficiency virus type 1]	0.008		
1		gi 6324561 ref NP_014630.1  required to degrade			
Į.	ļ	misfolded ER lumenal and integral membrane proteins;			
		Hrd1p [Saccharomyces cerevisiae] pir S66695 probable			
ì		membrane protein YOL013c - yeast (Saccharomyces			
l		cerevisiae) emb CAA99012.1  (Z74755) ORF YOL013c			
599	6324561	[Saccharomyces cerevisiae]	6.7		
1					
		gi 7494878 pir  T33047 hypothetical protein B0344.2 -			
ŀ		Caenorhabditis elegans gb AAC16982.1  (AF067209)			
600	7494878	B0344.2 gene product [Caenorhabditis elegans]	0.94		
Į					
		gi 11347199 pir  C81351 probable UDP-N-			
}		acetylmuramoylalanyl-D-glutamyl-2, 6-diaminopimelate-			
l		-D-alanyl-D-alanine ligase (EC 6.3.2.15) Cj0795c			
l	1	[imported] - Campylobacter jejuni (strain NCTC 11168)			
ļ		emb CAB73060.1  (AL139076) putative UDP-N-			
l		acetylmuramoylalanyl-D-glutamyl-2, 6-diaminopimelate			
601	11347199	- ligase [Campylobacter jejuni]	0.55		
}					
	1000-1-	gi 1293067 gb AAB06014.1  (U50958) drosophila seven-			
604	1293067	in-absentia gene product homolog [Mus musculus]	7.9		

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
		gi 5031975 ref NP 005875.1  p21-activated kinase 4;	
	ļ	protein kinase related to S. cerevisiae STE20, effector	
	l	for Cdc42Hs [Homo sapiens] ref[XP 041095.1] protein	
		kinase related to S. cerevisiae STE20, effector for	
		Cdc42Hs [Homo sapiens] ref[XP_041093.1  protein	
	1	kinase related to S. cerevisiae STE20, effector for	
		Cdc42Hs [Homo sapiens] sp O96013 PAK4_HUMAN	
٠.		SERINE/THREONINE-PROTEIN KINASE PAK 4	
	1	(P21-ACTIVATED KINASE 4) (PAK-4)	
	1	gb AAD01210.1  (AF005046) serine/threonine kinase	
606	5021075	[Homo sapiens] emb CAA09820.1  (AJ011855) PAK4	
606	3031975	protein [Homo sapiens]	1.1
608	0200205	gi 9280285 dbj BAB01686.1  (AB046104) unnamed protein product [Macaca fascicularis]	0.7
608	9280285		8.7
		gi 5869819 cmb CAB55576.1  (AJ249395) NADH- ubiquinoue oxidoreductase subunit 1 [Globodera	
609	5869819		0.04
609	2869819		0.84
Ì	1	gi 12045283 ref NP_073094.1  conserved hypothetical	
	l	protein [Mycoplasma genitalium]	
		sp P47662 Y423_MYCGE HYPOTHETICAL	
	ľ	PROTEIN MG423 pir  G64246 conserved hypothetical protein MG423 - Mycoplasma genitalium	
		gb AAC71647.1  (U39724) conserved hypothetical	
611	12045283	protein [Mycoplasma genitalium]	2.8
011	12043263	gi 11289932 pir  T50247 probable helicase [imported] -	2.6
		fission yeast (Schizosaccharomyces pombe)	
	l	emb CAB71840.1  (AL138666) putative helicase	
612	11289932	[Schizosaccharomyces pombe]	4.5
		gi 6119709 cmb CAB59566.1  (AJ249986) C3G protein	
613	6119709	[Rattus norvegicus]	1.6
		gi 7144507 gb AAA58585.2  (U12823) hemolysin	
620	7144507	[Acanthamoeba polyphaga]	8.1
		· · · · · · · · · · · · · · · · · · ·	
_		gi 15011757 gb AAB04582.3  (U64603) Hypothetical	
622	15011757	protein C09B7.1 [Caenorhabditis elegans]	7
		gi 8923324 ref NP 060247.1  hypothetical protein	
		FLJ20345 [Homo sapiens] ref[XP 008161.3]	
		hypothetical protein FLJ20345 [Homo sapiens]	
		dbj[BAA91105.1] (AK000352) unnamed protein product	
625	8923324	[Homo sapiens]	0.004

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO -	N	DESCRIPTION	P VALUE
		gi 6960319 gb AAD43326.2 AF155156_1 (AF155156)	
		adaptor-related protein complex AP-4 epsilon subunit	
626	6960319	[Homo sapiens]	0.00008
	1	gi 5739387 gb AAD50450.1 AF169388_1 (AF169388)	
628	5739387	alpha 4 collagen IV [Mus musculus]	5.5
		gi 3510234 gb AAC33487.1  (AC005581) R31237_1,	
631	3510234	partial CDS [Homo sapiens]	8.6
		gi 802150 gb AAB32775.1  (S75037) pancreatic	
		peptidylglycine alpha-amidating monooxygenase,	
		PAM=membrane-bound isoform {alternatively spliced,	
		clone PAM-3, transmembrane domain (Ba region)}	
		[human, islet cell tumor cell line QGP-1, Peptide Partial,	
632	802150	971 aa] [Homo sapiens]	1E-13
		gi 2105238 gb AAB57954.1  (U86905) similar to variola	
		and vaccinia E9L [Molluscum contagiosum virus	
636	2105238		1.5
		gi 7576305 emb CAB88003.1  (AJ277410) NADH	
641	7576305	dehydrogenase subunit 1 [Taenia hydatigena]	9.3
		gi 12231395 gb AAG49078.1 AF216121_1 (AF216121)	
643	12231395	phytochrome B [Gyminda tonduzii]	7
		gi 281178 pir  JQ1580 major surface antigen - hepatitis	
		B virus (subtype adw4q-, strains CNTS-38 and Fou)	
		emb CAA53344.1  (X75658) surface antigen [Hepatitis	
647	281178	B virus]	6.7
		gi 7503145 pir  T16315 hypothetical protein F41C3.6 -	
		Caenorhabditis elegans gb AAC46813.1  (U23521)	
		asked finisher to look for frameshift because of	
		discrepancy between prediction and EST CEESW76F;	
		looks like there should be a splice from 21751 to 21706.	
652	7502145	But no problem was found in the area [Caenorhabditis	
032	7503145	elegans]	6.9
		gi 14773502 ref XP_043252.1  PCAF associated factor	
		65 alpha [Homo sapiens] gb AAH08785.1 AAH08785 (BC008785) PCAF associated factor 65 alpha [Homo	
653	14773502		3E-10
022	11/13302	gi 14732840 ref XP_034110.1  hypothetical protein	3E-10
658	14732840	XP 034110 [Homo sapiens]	1.8
056	14732040	AT _024110 [rigino sapiens]	1.0
		gi 13442965 gb AAK26242.1 AF247132 1 (AF247132)	
659	13442965	putative chromatin remodeling factor [Mus musculus]	4
035	134472903	Perental curonistin remodering ractor [tyrus musculus]	4

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 4512671 gb AAD21725.1  (AC006931) En/Spm-like	
		transposon protein [Arabidopsis thaliana]	
		gb AAD33868.1 AF141375_1 (AF141375) protodermal	
		factor 1 [Arabidopsis thaliana]	
	i	gb AAD33869.1 AF141376_1 (AF141376) protodermal	
660	4512671	factor 1 [Arabidopsis thaliana]	6.6
		gi 13385330 ref[NP_080127.1  RIKEN cDNA	
		1700010I14 gene [Mus musculus] dbj BAB24262.1	
663	13385330	(AK005830) putative [Mus musculus]	0.33
		gi 14325583 dbj BAB60486.1  (AP000996) unknown	
665	14325583	product [Thermoplasma volcanium]	5.4
		gi 14779941 ref XP_007847.3  hypothetical protein from	
666	14779941	clone 24796 [Homo sapiens]	0.00000002
		gi 14330407 emb CAC41079.1  (AJ308518) P2X5	
670	14330407	receptor [Gallus gallus]	3,3
		gi 13991595 gb AAK51427.1  (AF355796) BRCA1	
672	13991595	[Echymipera kalubu]	6.9
		gi 14728540 ref XP_048261.1  similar to ring finger	
		protein 23; RING-B box-coiled coil-B30.2 (M.	
681	14728540	musculus) [Homo sapiens]	0.026
		gi 14740222 ref XP 049918.1  similar to hypothetical	
683	14740222	protein MGC10940 (H. sapiens) [Homo sapiens]	1E-86
- 005	11710222	gi 11359451 pir  T51040 hypothetical protein	12-00
		B15I20.100 [imported] - Neurospora crassa	
		emb CAB97464.1  (AL389900) conserved hypothetical	
684	11359451	protein [Neurospora crassa]	9.2
		gi 14735526 ref XP 001804.4  guanylate binding protein	
685	14735526	2, interferon-inducible [Homo sapiens]	2.3
		, , , , , , , , , , , , , , , , , , , ,	
		gi 7510353 pir T27275 hypothetical protein Y63D3A.9	
		Caenorhabditis elegans emb CAA21711.1  (AL032652)	
		contains similarity to Pfam domain: PF00646 (F-box	
	1	domain.), Score=43.2, E-value=1.9e-09, N=1	
687	7510353	[Caenorhabditis elegans]	6.8
		gi 14773502 ref XP 043252.1  PCAF associated factor	
		65 alpha [Homo sapiens] gb AAH08785.1 AAH08785	
		(BC008785) PCAF associated factor 65 alpha [Homo	
697	14773502		3E-10
		gi 7498998 pir  T16057 hypothetical protein F13D11.2 -	
700	7498998	Caenorhabditis elegans	0.27

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
1	ŀ	744000001 1101700000 11417004550	
		gi 4493893 emb CAB39002.1  (AL034558) predicted	
703	4493893	using hexExon; MAL3P2.15 (PFC0230c), Hypothetical protein len: 3979 aa [Plasmodium falciparum]	0.2
703	4493693	gi/7243081 dbi BAA92588.1  (AB037771) KIAA1350	9.2
706	7243081	protein [Homo sapiens]	1E-79
		gi 7496384 pir  T33266 hypothetical protein C24B9.10 -	XD
l		Caenorhabditis elegans gb[AAC19256,1] (AF068709)	
		Hypothetical protein C24B9.10 [Caenorhabditis	
707	7496384	elegans]	5.4
		gi 7492007 pir  T41680 hypothetical protein	
l		SPCP1E11.01c - fission yeast (Schizosaccharomyces	
		pombe) emb CAB54860.1  (AL117183) hypothetical	
708	7492007	protein [Schizosaccharomyces pombe]	4.1
	l <b>_</b>	gi 14758507 ref XP_044768.1  similar to death receptor	
709	14758507	6 (H. sapiens) [Homo sapiens]	0.031
İ		-11/25/12/25/1/14 A FT 149/20 1/4 CO11 CC4 11 (4 CO11 CC4)	
710	4512025	gi 6513925 gb AAF14829.1 AC011664_11 (AC011664) unknown protein [Arabidopsis thaliana]	0.83
710	0313923	unknown protein [Arabidopsis manana]	0.83
		gi 6434714 emb CAB61161.1  (AL132973) putative	
713	6434714	membrane protein. [Streptomyces coelicolor A3(2)]	1.2
		[	
į .		gi 8810476 gb AAF80137.1 AC024174 19 (AC024174)	
		Contains similarity to a hypothetical protein F24K9.13	
ŀ		gi 6006885 from Arabidopsis thaliana gb AC008153	
714	8810476	and contains multiple PPR PF 01535 repeats	0.22
		gi 9626839 ref[NP_041109.1  ORF 18 [ictalurid	
		herpesvirus 1] sp Q00120 VG18_HSVI1	
1		HYPOTHETICAL GENE 18 PROTEIN pir A36788	
		hypothetical protein ORF18 - ictalurid herpesvirus 1	
710	0606000	(strain auburn 1) gb AAA88121.1  (M75136) ORF 18	
719	9626839	[ictalurid herpesvirus 1]	9.2
		gi 15004903 gb AAK77203.1 AC006622_3	
723	15004903	(AC006622) Hypothetical protein C52D10.12 [Cacnorhabditis elegans]	2.4
123	13004903	[Cachornaoun's cicgans]	2,4
		gi 6677663 ref[NP 033047.1  retinoic acid induced 1	
1		Mus musculus] pir T30250 GT1 protein - mouse	
1		dbj[BAA06184.1] (D29801) Unknown [Mus musculus]	
724	6677663	prf[2123391A GT1 gene [Mus musculus]	0.12

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 2343183 gb AAB67715.1  (AF013243) zinc finger	
729	2343183	transcription factor [Lytechinus variegatus]	0.021
		gi 7465334 pir  B71980 proline dehydrogenase (EC	
1		1.5.99.8) / 1-pyrroline-5-carboxylate dehydrogenase	
		(EC 1.5.1.12) putA [similarity] - Helicobacter pylori	
)	l	(strain J99) gb AAD05632.1  (AE001444)	
		Proline/pyrroline-5-carboxylate dehydrogenase	
730	7465334	[Helicobacter pylori J99]	4.9
		gi 9631436 ref NP_048287.1  ORF MSV216 SCG gene	
		family protein [Melanoplus sanguinipes	
		entomopoxvirus] pir  T28377 ORF MSV216 SCG gene	
}		family protein - Melanoplus sanguinipes	
ì		entomopoxvirus gb AAC97749.1  (AF063866) ORF	
		MSV216 SCG gene family protein [Melanoplus	
733	9631436	sanguinipes entomopoxvirus]	0.52
l		gi 13813168 gb AAK40402.1  (AE006645) Hypothetical	
734	13813168	protein [Sulfolobus solfataricus]	2.7
		gi 14776978 ref XP_033086.1  hypothetical protein	
736	14776978	XP_033086 [Homo sapiens]	5.4
		gi 7294863 gb AAF50194.1  (AE003550) CG6718 gene	
737	7294863	product [Drosophila melanogaster]	1.9
		gi 7465334 pir  B71980 proline dehydrogenase (EC	
		1.5.99.8) / 1-pyrroline-5-carboxylate dehydrogenase	
1	<b>\</b>	(EC 1.5.1.12) putA [similarity] - Helicobacter pylori	
		(strain J99) gb AAD05632.1  (AE001444)	
745	7465224	Proline/pyrroline-5-carboxylate dehydrogenase	2.0
743	7405334	[Helicobacter pylori J99]	3.9
ĺ		-3575021(1-114 AD51020 114 P172200 1 (4 P172200)	
		gi 5759216 gb AAD51030.1 AF172399_1 (AF172399) p75 neurotrophin receptor a-1 [Xenopus laevis]	
1		gb AAD51031.1 AF172400 1 (AF172400) p75	
750	5759216	neurotrophin receptor a-2 [Xenopus laevis]	3.9
150	3133210	gi 6518511 dbi BAA87907.1  (AB021222) ketosynthase	3.5
751	6518511	[Streptomyces rochei]	6.6
		L	
		gi 6679777 ref NP 032029.1  fibroblast growth factor	
1	1	15 [Mus musculus] sp O35622 FGFF MOUSE	
		FIBROBLAST GROWTH FACTOR-15	
1		PRECURSOR (FGF-15) gb AAB63197.1  (AF007268)	
1		fibroblast growth factor [Mus musculus]	
754	6679777	dbj BAB30961.1  (AK017829) putative [Mus musculus]	7.7

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS			
NO	N	DESCRIPTION	P VALUE	
		THE ACCOUNT OF THE PARTY OF THE		
ł	i	gi 7498881 pir  T20748 hypothetical protein F11A5.2 -		
		Caenorhabditis elegans emb CAB07355.1  (Z92830) contains similarity to Pfam domain: PF01604 (TTM		
	ļ.	chemoreceptor), Score=428.7, E-value=1.7e-125, N=1		
755	7498881	[Caenorhabditis elegans]	6.4	
133	7420001	gi 1378135 gb AAB02606.1  (U58466) putative	0.4	
1	ì	fibroblast growth factor receptor 1-like protein; one of		
		three possible open reading frames in the sequence		
756	1378135	[Rattus norvegicus]	5.1	
750	1376133	gi 7444901 pir  B71630 hypothetical protein RP702 -	3.1	
	l	Rickettsia prowazekii emb CAA15138.1  (AJ235273)		
758	7444901	unknown [Rickettsia prowazekii]	2.2	
/50	744701	gil12854427 dbi BAB30025.1  (AK015906) putative	2.2	
759	12854427		6.6	
707	12001127	gi 4185892 emb CAA21831.1  (AL033125) 1-	0.0	
		evidence=predicted by content~1-		
	i	method=genefinder;084~1-evidence end~2-		
	Į.	evidence=predicted by match~2-		
		match accession=SWISS-PROT:P38205~2-		
		match description=HYPOTHETICAL 77.9 KD		
	- 33	PROTEIN IN RRN10-MCM2 INTERGENIC		
763	4185892		9.1	
	1	gi 14725855 ref XP 038865.1  similar to hypothetical		
766	14725855		9.2	
		gi 7582302 gb AAF64271.1 AF208857 1 (AF208857)		
770	7582302	BM-015 [Homo sapiens]	8E-13	
		gi 6594283 dbj BAA88419.1  (AB016615) hydrophobic		
772	6594283	transmembrane protein [Staphylococcus aureus]	1.7	
		gi 12597803 gb AAG60115.1 AC073178_26		
783	12597803	(AC073178) hypothetical protein [Arabidopsis thaliana]	0.62	
		gi 8777303 dbj BAA96893.1  (AB018112) serine		
784	8777303	carboxypeptidase [Arabidopsis thaliana]	1.2	
		gi 10045258 emb CAC07930.1  (AJ228487) cytochrome		
785	10045258	c oxidase subunit 3 [Acanthogammarus godlewskii]	1.5	
		gi 1794167 dbj BAA11217.1  (D78137) unnamed		
786	1794167	protein product [Vibrio parahaemolyticus]	9.7	
		gi 12835698 dbj BAB23329.1  (AK004489) putative		
790	12835698	[Mus musculus]	8.1	

	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Pr			
SEQ ID	ACCESS			
NO	N	DESCRIPTION	P VALUE	
		gi 131761 sp P11636 QAY_NEUCR QUINATE		
		PERMEASE (QUINATE TRANSPORTER)		
i		pir  G31277 quinate transport protein - Neurospora		
		crassa (tentative sequence) emb CAA32752.1  (X14603)		
791	131761	quinate transporter [Neurospora crassa]	1.7	
1	i .	gi 11640576 gb AAG39285.1 AF113214_1 (AF113214)		
794	11640576	MSTP034 [Homo sapiens]	6.3	
		gi 5306243 gb AAD41976.1 AC006438_8 (AC006438)		
797	5306243	hypothetical protein [Arabidopsis thaliana]	3.4	
		Umanaged : Impenses 1.1		
		gi 7511787 pir  T29089 alpha-mannosidase (EC 3.2.1),		
798	7511787	class II - fall armyworm gb AAB62719.1  (AF005034)	7	
198	/311/6/	alpha-mannosidase II [Spodoptera frugiperda]		
		gi 3170793 gb AAC18201.1  (AF062165) immunoglobulin heavy chain variable region [Homo		
800	3170793	sapiens]	8.7	
800	3170793		8.7	
		gi 11466458 ref NP_038161.1  NADH dehydrogenase		
		subunit 4L [Chrysodidymus synuroideus]		
		gb AAB95105.1  (U54633) NADH dehydrogenase subunit 4L [Chrysodidymus synuroideus]		
1	1	gblAAF36927.1lAF222718 1 (AF222718) NADH		
		dehydrogenase subunit 4L [Chrysodidymus		
802	11466458	synuroideus]	4.2	
- 002	11100130	gi 14779941 ref XP 007847.3  hypothetical protein from	7.2	
803	14779941	clone 24796 [Homo sapiens]	0.00000002	
	11////	gi 13430868 ref NP 077008.1  hypothetical protein	0.0000002	
1		MGC2615 [Homo sapiens] ref[XP_015759.1]		
		hypothetical protein MGC2615 [Homo sapiens]		
		gb AAH01656,1 AAH01656 (BC001656) Unknown		
804	13430868	(protein for MGC:2615) [Homo sapiens]	4E-21	
		gi 3293234 gb AAC25913.1  (U96413) T-DNA		
808	3293234	oncoprotein [Agrobacterium tumefaciens]	7.9	
		gi 7299169 gb AAF54367.1  (AE003682) Fps85D gene		
810	7299169	product [alt 1] [Drosophila melanogaster]	5.6	
		gi 11361627 pir  H82736 hypothetical protein XF0992		
		[imported] - Xylella fastidiosa (strain 9a5c)		
		gb AAF83802.1 AE003937_6 (AE003937) hypothetical		
814	11361627	protein [Xylella fastidiosa 9a5c]	0.43	
		gi 10803160 emb CAC13088.1  (AL445503) putative		
		two component system response regulator [Streptomyces		
816	10803160	coelicolor]	6.4	

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PCT/US01/25840

WO0214500 [fle //E /WO0214500 opc]

WO 02/14500

#### Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins) SEO ID ACCESS NO Ν DESCRIPTION P VALUE gi|2613090|gb|AAB84279.1| (AF030317) OmpR 819 2613090 [Proteus vulgaris] 1.7 gil2494232|splO62814|E2F5 RAT TRANSCRIPTION FACTOR E2F5 (E2F-5) gb|AAB00180.1| (U31668) 821 2494232 E2F-5 [Rattus norvegicus] 7.8 gil14729752|reflXP 036388.1| hypothetical protein 822 14729752 MGC5297 [Homo sapiens] 6.7 gi|7024427|emb|CAA87594.2| (Z47547) Pwi=orf234.1;Mpo=orf244;Angio=orfx~no ATG start 825 7024427 codon [Chondrus crispus] 8 gi|11691811|emb|CAC11114.1| (AL121893) bA189K21.6 (Sec23 (S. cerevisiae) homolog B) [Homo 833 11691811 sapiens] 5.6 gi|117704|sp|P18246|CXA1 BOVIN GAP JUNCTION ALPHA-1 PROTEIN (CONNEXIN 43) (CX43) (VASCULAR SMOOTH MUSCLE CONNEXIN 43) pir||A36623 gap junction protein Cx43 - bovine gb|AAA30459.1| (J05535) vascular smooth muscle 834 117704 connexin43 [Bos taurus] 9.2 gil1170606lsplP43188lKADC MAIZE ADENYLATE KINASE, CHLOROPLAST (ATP-AMP TRANSPHOSPHORYLASE) pirl|\$45634 adenylate kinase (EC 2.7.4.3), chloroplast - maize pdb|1ZAK|A Chain A. Adenvlate Kinase From Maize In Complex With The Inhibitor P1, P5-Bis(Adenosine-5'-)pentaphosphate (Ap5a) pdb|1ZAK|B Chain B, Adenylate Kinase From Maize In Complex With The Inhibitor P1.P5-Bis(Adenosine-5'-)pentaphosphate 835 1170606 (Ap5a) 0.0000001 gil7661750|refINP 054866.1| HSPC047 protein [Homo sapiens] gblAAF29019.1lAF161532 1 (AF161532) 836 7661750 HSPC047 [Homo sapiens] 5E-28 gi|13815530|gb|AAK42398.1| (AE006828) Transport 13815530 protein, hypothetical [Sulfolobus solfataricus] 838 5 gi|14771844|ref|XP 045585.1| hypothetical protein FLJ11085 [Homo sapiens] emb[CAB91047.2] 839 14771844 (AL109935) dJ1022P6.2 (KIAA1434) [Homo sapiens] 0.089 gil4007786[emblCAA51374.1] (X72850) acetyl CoA 843 4007786 acetyltransferase [Sphingomonas sp.] 4.6

	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESS				
NO	N	DESCRIPTION	P VALUE		
		gi 9968471 emb CAC06698.1  (AJ275988) Kruppel like			
847	9968471	factor [Mus musculus]	6.2		
		gi 7499805 pir  T21337 hypothetical protein F25D7.1 -			
		Caenorhabditis elegans emb CAB01696.1  (Z78418)			
		cDNA EST yk60g6.3 comes from this gene~cDNA EST			
		yk60g6.5 comes from this gene~cDNA EST			
		yk156b10.3 comes from this gene~cDNA EST			
		yk156b10.5 comes from this gene~cDNA EST			
848	7499805	yk399f2.3 comes from this gene~cDNA EST yk399f2.5			
848	7499803	comes from this gene~cDN>	7.1		
		gi 6581093 gb AAF18453.1 AF205599 1 (AF205599)			
849	6581093	transposase-like protein [Mus musculus]	4.4		
042	0301073	transposase-nac protein [ivids musculus]	4.4		
		gi 13651256 ref[XP_015436.1  chloride channel 4			
		[Homo sapiens] ref[XP 045758.1] chloride channel 4			
		[Homo sapiens] ref[XP_045756.1] chloride channel 4			
]		[Homo sapiens] ref[XP_045757.1] chloride channel 4			
		[Homo sapiens] dbj BAA77327.1  (AB019432) chloride			
		channel protein 4 [Homo sapiens]			
		gb AAD50981.1 AF170492_1 (AF170492) chloride			
853	13651256	channel CLC4 [Homo sapiens]	2.3		
		gi 336159 gb AAA46774.1  (M90520) polymerase			
854	336159	protein [Woodchuck hepatitis B virus]	3.2		
		gi 8979124 dbj BAA98959.1  (AP002547) CT651			
861	8979124	hypothetical protein [Chlamydophila pneumoniae J138]	0.76		
i					
		gi 4758712 ref NP_004659.1  alpha-glucosidase; brush			
	ĺ	border hydrolase [Homo sapiens]			
		sp O43451 MGA_HUMAN MALTASE-			
		GLUCOAMYLASE, INTESTINAL [INCLUDES:			
		MALTASE (ALPHA-GLUCOSIDASE); GLUCOAMYLASE (GLUCAN 1,4-ALPHA-			
l		GLUCOSIDASE)] gb AAC39568.1  (AF016833)			
863	4758712	maltase-glucoamylase [Homo sapiens]	0.1		
- 505	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	gi 13812050 ref NP 113184.1  hypothetical protein	U.1		
		[Guillardia thetal gblAAK39752.1 AF083031 109			
864	13812050	(AF083031) hypothetical protein [Guillardia theta]	4.3		
		gi 10726408 gb AAF54394.2  (AE003683) CG9381			
867	10726408	gene product [Drosophila melanogaster]	5.5		
867	10726408	gene product [Drosophila melanogaster]	5.5		

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 7299899 gb AAF55074.1  (AE003705) CG7987 gene	
		product [alt 1] [Drosophila melanogaster]	
		gb AAF55075.1  (AE003705) CG7987 gene product [alt	
870	7299899	2] [Drosophila melanogaster]	9.7
	}	-ilcozoczal-bia Arcadalo Illa Euganoa G (A Euganoa)	
972	(070225	gi 6979325 gb AAF34418.1 AF172282_7 (AF172282)	4.0
872	6979325	putative phosphatidylinositol 4-kinase [Oryza sativa]	4.8
		gi 13699930 dbj BAB41229.1  (AP003129) conserved	
	}	hypothetical protein [Staphylococcus aurcus subsp.	
		aureus N315] dbj BAB56175.1  (AP003358) conserved	
873	12600020	hypothetical protein [Staphylococcus aureus subsp. aureus Mu50]	2.7
8/3	13099930	gi 6841256 gb AAF28981.1 AF161421 1 (AF161421)	3.7
877	6841256		777.64
8//	0041230	gi 2459878 gb AAC40459.1  (AF005734) glycoprotein	7E-64
879	2459878	precursor [Marburg virus]	2.5
8/3	2439076	gil14759292 refiXP 006700,2  checkpoint with forkhead	2.3
		and ring finger domains [Homo sapiens]	
	ì	gblAAF91084.1lAF170724 1 (AF170724) cell cycle	
880	14759292		0.000000002
880	14737272	checkpoint protein CTI K [Fromo sapiens]	0,00000002
	ì	gi 141164 sp P19297 YORM TTV1 HYPOTHETICAL	
	1	38.6 KD PROTEIN emb CAA32993.1  (X14855) URF	
886	141164	(352 AA) [Thermoproteus tenax virus 1]	2.2
	111101	gi 1755061 gb AAB50394.1  (U62737) photosystem I	
889	1755061	subunit XI [Synechococcus sp. PCC 7942]	3.6
		gi 7464739 pir  G71920 hypothetical protein jhp0518 -	
	Į.	Helicobacter pylori (strain J99) gb AAD06099.1	
890	7464739	(AE001485) putative [Helicobacter pylori J99]	7.6
		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	
	1	gi 14141161 reffNP 004492.2  heterogeneous nuclear	
	1	ribonucleoprotein U, isoform b; hnRNP U protein;	
	l	scaffold attachment factor A; p120 nuclear protein	
		[Homo sapiens] gb AAH03367.1 AAH03367	
		(BC003367) heterogeneous nuclear ribonucleoprotein U	
	}	(scaffold attachment factor A) [Homo sapiens]	
l	1	gb AAH03621.1 AAH03621 (BC003621) heterogeneous	
		nuclear ribonucleoprotein U (scaffold attachment factor	
891	14141161	A) [Homo sapiens]	6.9
		gi 7649887 dbj BAA94165.1  (AP000422) tail fiber	
892	7649887	protein [Escherichia coli O157:H7]	0.63

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		Toble 3D Nessest Naidle - (Dl - 4V - N - D - 1 - 1 - 1	P. dia
SEQ ID	ACCESS	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
NO	N N	DESCRIPTION	P VALUE
		gi 7496871 pir  T19650 hypothetical protein C32H11.7 -	
	ļ	Caenorhabditis elegans emb CAB05141.1  (Z82260)	
		cDNA EST yk576f6.3 comes from this gene	
901	7496871	[Cacnorhabditis elegans]	4.6
005	10400100	gi 13400109 gb AAK21974.1  (U77931) rRNA	
905	13400109	promoter binding protein [Rattus norvegicus]	0.003
918	12225450	gi 13235458 cmb CAC33751.1  (AJ293322) ProP2	
918	13233438	protein [Rickettsia typhi]	4.1
919	7202165	gi 7292165 gb AAF47577.1  (AE003472) CG12023 gene product [Drosophila melanogaster]	0.25
919	7292165		0.25
		gi 12831207 ref NP_075579.1  gamma-aminobutyric	
		acid (GABA-A) receptor, subunit epsilon [Rattus	
	1	norvegicus] gb AAF70383.1 AF189262_1 (AF189262)	
920	10021007	GABA-A receptor epsilon-like subunit [Rattus	4.5
920	12831207	norvegicus]	4,5
		gi 10765285 gb AAG22971.1 AF183431 1 (AF183431)	
926	10765205	inhibitor of apoptosis protein 2 [Rattus norvegicus]	4.2
920	10703263	gi 2832268 gb AAC15477.1  (AF043233) Caco-2	4.2
927	2832268	oligopeptide transporter [Homo sapiens]	5.1
921	2032200	ongopeptide transporter [riomo sapiens]	3.1
		gi 6179898 gb AAF05702.1 AF190129 1 (AF190129)	
928	6170909	Na+/K+/2Cl- cotransporter [Callinectes sapidus]	6.1
920	01/9090		0.1
	ĺ	gi 12644035 sp Q9Z8M1 IF2_CHLPN	
	1	TRANSLATION INITIATION FACTOR IF-2	
		pir  E81576 translation initiation factor 2 CP0440	
	ì	[imported] - Chlamydophila pneumoniae (strain AR39) gb AAF38279.1  (AE002205) translation initiation	
		factor 2 [Chlamydophila pneumoniae AR39]	
	1	dbj BAA98527.1  (AP002546) initiation factor-2	
931	12644035	[Chlamydophila pneumoniae J138]	0.91
931	12044033	[Cinantydopinia pileuniomae 1138]	0.91
		"11005C3C3  (BVB 052011 11 3201 115 ED '31	
		gi 10956362 ref[NP_052811.1  pXO1-115 [Bacillus anthracis] pir  C59105 hypothetical protein pXO1-115 -	
		Bacillus anthracis virulence plasmid pXO1	
		gb AAD32419.1 AAD32419 (AF065404) pXO1-115	
		[Bacillus anthracis] gblAAD41347.1[AF150965 1	
932	10956362	(AF150965) resolvase X [Bacillus anthracis]	1.9
732	10550502	(THE TOO SO TO SO TRUSCO TE [DECINGS ANGUIACIS]	1,3
		gi 13814214 gb AAK41294.1  (AE006722) Conserved	
933	13814214	hypothetical protein [Sulfolobus solfataricus]	0.99
,,,,	13014214	nypoulousus protein (puriototus sotiataricus)	0,55

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 6320195 ref NP_010275.1  Ydl009cp	
		[Saccharomyces cerevisiae] pir  S52510 probable	
	İ	membrane protein YDL009c - yeast (Saccharomyces	
		cerevisiae) emb CAA88350.1  (Z48432) unknown	
		[Saccharomyces cerevisiae] emb CAA98568.1	
940	6320195	(Z74059) ORF YDL009c [Saccharomyces cerevisiae]	7
		gi 10181130 ref[NP_065586.1  acid sphingomyelinase-	
		like phosphodiesterase 3a [Mus musculus]	
942	10101120	emb CAA69329.1  (Y08135) acid sphingomyelinase-like	CT 04
942	10181130	phosphodiesterase [Mus musculus]	6E-84
		gi 13124718 sp P54358 DPOD_DROME DNA	
		POLYMERASE DELTA CATALYTIC SUBUNIT	
943	12124710	gb AAF49555.1  (AE003529) DNApol-delta gene product [Drosophila melanogaster]	2.8
943	13124/18	gi 2258300 gb AAB63269.1  (AF001783) AgrC	2.8
946	2258300	[Staphylococcus aureus]	1.9
940	2238300	[Staphylococcus aureus]	1.9
		gi 7460037 pir  T13431 hypothetical protein T17A13.20	
		Arabidopsis thaliana emb CAB79678.1  (AL161574)	
947	7460037	hypothetical protein [Arabidopsis thaliana]	3.4
	7100007	gi 7494291 pir  E71616 hypothetical protein PFB0365w	
		malaria parasite (Plasmodium falciparum)	
		gb AAC71862.1  (AE001390) hypothetical protein	
948	7494291	[Plasmodium falciparum]	1.9
		gi 1707627 emb CAA96381.1  (Z71701) cytochrome	
950	1707627	oxidase subunit III [Euhadra herklotsi]	9.1
		gi 6739602 gb AAF27330.1  (AF178534) talin [Homo	
952	6739602	sapiens]	1E-14

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	Description.	
NO	N	DESCRIPTION	P VALUE
		gi 7498956 pir  T20804 hypothetical protein F12F6.1 -	
		Caenorhabditis elegans emb CAA97791.1  (Z73425)	
	1	cDNA EST yk30a8.3 comes from this gene~cDNA EST	
		yk12d11.5 comes from this gene~cDNA EST yk30a8.5	
		comes from this gene~cDNA EST yk117g9.3 comes from this gene~cDNA EST yk113h11.5 comes from this	
	Ì	gene~cDNA EST yk117g9.5 comes from this	
		gene~cDNA> emb CAA97796.1  (Z73426) cDNA EST	
		yk30a8.3 comes from this gene~cDNA EST yk12d11.5	
		comes from this gene~cDNA EST vk30a8.5 comes from	
		this gene~cDNA EST yk117g9.3 comes from this	
		gene~cDNA EST yk113h11.5 comes from this	
		gene~cDNA EST yk117g9.5 comes from this	
954	7498956	gene~cDNA>	6.6
1		gi 9631417 ref NP 048325.1  ORF MSV254 leucine	
[		rich repeat gene family protein, similar to Amsacta	
ĺ		moorei entomopoxvirus Q3 ORF SW:P28854	
		[Melanoplus sanguinipes entomopoxvirus] pir T28415	
ļ	,	ORF MSV254 leucine rich repeat gene family protein -	,
l		Melanoplus sanguinipes entomopoxvirus	
		gb AAC97730.1  (AF063866) ORF MSV254 leucine	
		rich repeat gene family protein, similar to Amsacta	
		moorei entomopoxvirus Q3 ORF SW:P28854	
955	9631417	[Melanoplus sanguinipes entomopoxvirus]	2.1
		gi 7160126 emb CAB76312.1  (AL158060) putative	
956	7160126	membrane protein. [Streptomyces coclicolor A3(2)]	0.25
050	1202065	gi 1203965 gb AAA89173.1  (LA2379) bone-derived	l ,,
959	1203965	growth factor [Homo sapiens]	1.4
		cils 95 147lociD29004IELLIC EMENI ELLIC PROTEINI	
		gi 585147 sp P38094 FLUG_EMENI FLUG PROTEIN pir  A53186 fluG protein - Emericella nidulans	
960	585147	gb AAC37414.1  (L27817) FluG [Emericella nidulans]	5.4
700	303147		J. <del>4</del>
		gi 11352695 pir  D83270 transcription-repair coupling protein Mfd PA3002 [imported] - Pscudomonas	
		aeruginosa (strain PAO1)	
		gb AAG06390,1 AE004725 3 (AE004725)	
		transcription-repair coupling protein Mfd [Pseudomonas	
963	11352695	aeruginosa]	1.3
			L—

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 119777 sp P16291 FA9 SHEEP COAGULATION	
		FACTOR IX (CHRISTMAS FACTOR) pir   147078	
		coagulation factor IXa (EC 3.4.21.22) - sheep	
		(fragment) gb AAA31520.1  (M26233) factor IX [Ovis	
964	119777	aries]	8.2
			-
	ł	gi 7473507 pir  F75587 probable glycosyltransferase -	
	ł	Deinococcus radiodurans (strain R1)	
	ĺ	gb AAF12451.1 AE001863_76 (AE001863)	
965	7473507	glycosyltransferase, putative [Deinococcus radiodurans]	0.24
		gi 2494130 gb AAB80639.1  (AC002376) Contains	
	ŀ	similarity to Glycine SRC2 (gb AB000130).	
967	2494130	[Arabidopsis thaliana]	5.2
		gi 10728660 gb AAF52603.2  (AE003620) CG8683	
972	10728660	gene product [Drosophila melanogaster]	6E-12
		gi 7500037 pir  T34063 chromosome segregation protein	
		smc1 F28B3.7 [similarity] - Caenorhabditis elegans	
		gb AAK21378.1  (AF003136) contains similarity to	
980	7500037	ATP synthase subunit B [Caenorhabditis elegans]	0.68
		gi 7298917 gb AAF54122.1  (AE003675) CG10272	
981	7298917	gene product [Drosophila melanogaster]	5.9
		gi 1170115 sp P46430 GTT1_MANSE	
		GLUTATHIONE S-TRANSFERASE 1 (GST CLASS-	
		THETA) gb AAA92880.1  (L32091) glutathione S-	
983	1170115	transferase [Manduca sexta]	7.1
		gi 10864490 gb AAG24203.1  (AF022981) Hypothetical	
985	10864490	protein W03F9.6 [Caenorhabditis elegans]	5.3
		gi 105400 pir  A35648 B-cell adhesion protein CD22	
		alpha splice form precursor - human emb CAA36988.1	
987	105400	(X52785) CD22 antigen [Homo sapiens] prf  1608208A	
981	105400	B cell antigen DC22 [Homo sapiens]	6
989	14750376	gi 14750376 ref XP_012394.3  myosin IXA [Homo	3.9
969	14/303/6		3.9
		gi 6324560 ref NP_014629.1  Yol013w-ap	
		[Saccharomyces cerevisiae] pir  S78736 protein	
990	6324560	YOL013w-a - yeast (Saccharomyces cerevisiae) delta remnant	0.2
990	0324360	gi 7290503 gb AAF45956.1  (AE003431) CG3527 gene	9.3
991	7290503	gi[7290503[gb]AAF45956.1] (AE003431) CG3527 gene product [Drosophila melanogaster]	7.6
991	1290503	product [Drosopmia meianogaster]	7.6

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 9964623 ref NP_064753.1  RP DNA pol [Roseophage	
		SIO1] gb AAG02598.1 AF189021_17 (AF189021)	
994	9964623	Roseophage SIO1 complete genome	3.5
		gi 2769538 emb CAA11217.1  (AJ223279) voltage-	
996	2769538	sensitive sodium channel [Plutella xylostella]	7.6
ļ		gi 9628506 ref[NP_043384.1  regulatory protein E2	
		[Human papillomavirus type 29]	
		sp P50772 VE2_HPV29 REGULATORY PROTEIN	
		E2 gb AAA79432.1  (U31784) regulatory protein E2	
998	9628506		6.2
		gi 6566147 dbj BAA04745.2  (D21203) large Forked	
999	6566147	protein [Drosophila melanogaster]	0.085
		gi 11346920 pir  H81390 probable integral membrane	
		protein Cj0461c [imported] - Campylobacter jejuni	
		(strain NCTC 11168) cmb CAB75099.1  (AL139075)	
		putative integral membrane protein [Campylobacter	
1002	11346920	jejuni]	1.5
		gi 9294038 dbj BAB01995.1  (AB020746)	
		gene_id:MOB24.1~unknown protein [Arabidopsis	
1005	9294038	thaliana]	9.8
		gi 4505067 ref NP '002349.1  MAD2-like 1; mitotic	
		arrest deficient, yeast, homolog-like 1 [Homo sapiens]	
		sp Q13257 MD21 HUMAN MITOTIC SPINDLE	
		ASSEMBLY CHECKPOINT PROTEIN MAD2A	
		(MAD2-LIKE 1) (HSMAD2) pir  G01942 mitotic	
		feedback control protein Madp2 homolog - human	
		gb AAC52060.1  (U31278) mitotic feedback control	
ĺ		protein Madp2 homolog [Homo sapiens]	
		gb AAC50781.1  (U65410) Mad2 [Homo sapiens]	
		emb CAA03943.1  (AJ000186) MAD2 [Homo sapiens]	
		gb AAH00356.1 AAH00356 (BC000356) MAD2	
l		(mitotic arrest deficient, yeast, homolog)-like 1 [Homo	
		sapiens] gb AAH05945.1 AAH05945 (BC005945)	
l		MAD2 (mitotic arrest deficient, yeast, homolog)-like 1	
		[Homo sapiens] gb AAK38174.1  (AF202273) MAD2-	
1006	4505067	like protein 1 [Homo sapiens]	0.000005

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	DA/ALTIE
NO	IN IN		P VALUE
	1	gi 13994140 ref NP_038928.1  antigen p97 (melanoma	
ĺ		associated) identified by monoclonal antibodies 133.2	
	1	and 96.5 [Mus musculus] dbj BAA86655.1	
		(AB024336) membrane-bound transferrin-like protein	
	1	p97 [Mus musculus] dbj BAB41139.1  (AB047799)	
1010	12004140	membrane-bound transferrin-like protein [Mus	4.
1010	13994140	musculus]	4.1
		gi 11465559 ref NP 045049.1  unknown [Cyanidium	
l	ļ	caldarium] gb AAF12997.1 AF022186 169	
1011	11465559	(AF022186) unknown [Cyanidium caldarium]	1.8
		gi 13421288 gb AAK22158.1  (AE005691) TonB-	
1012	13421288	dependent receptor [Caulobacter crescentus]	0.65
l			
l		gi 6978481 ref NP_036905.1  a-kinase anchoring protein	
		[Rattus norvegicus] sp Q62924 AK11_RAT A KINASE	
		ANCHOR PROTEIN 11 (PROTEIN KINASE A	
		ANCHORING PROTEIN 11) (PRKA11) (A KINASE	
1	ŀ	ANCHOR PROTEIN 220 KDA) (AKAP 220)	
		pir  T42732 A-kinase anchoring protein AKAP 220 - rat	
l		gb AAB06559.1  (U48288) AKAP 220 [Rattus	
1013	6978481	norvegicus]	3
		gi 7498137 pir  T31694 hypothetical protein D1065.2 -	
1015	7498137	Caenorhabditis elegans	7.4
İ		gi 9944230 emb CAC05416.1  (AJ400866) membrane	
1017	9944230	tyrosine phosphatase [Bos indicus]	7
		gi 14520862 ref[NP_126337.1  hypothetical protein	
		[Pyrococcus abyssi] pir  G75106 hypothetical protein	-0.1
1		PAB0444 - Pyrococcus abyssi (strain Orsay)	
		emb CAB49568.1  (AJ248285) hypothetical protein	
1023	14520862	[Pyrococcus abyssi]	9.7
		1112003471 : HD02020	
1	1	gi 11290247 pir  D82039 conserved hypothetical protein	
		VC2740 [imported] - Vibrio cholerae (group O1 strain	
1025	11200247	N16961) gb AAF95879.1  (AE004339) conserved	0.2
1025	11290247	hypothetical protein [Vibrio cholerae] gi 12484269 gb AAG54048,1 AF224607 4 (AF224607)	8.2
		NADH dehydrogenase subunit 4 [Propithecus verreauxi	
1027	12484269		7
1027	12404209	gi 14329703 emb CAC40662.1  (AJ292926) anaerobic	<del>'</del>
i		gi 14329703 emb CAC40662.1  (AJ292926) anaerobic (class III) ribonucleotide reductase large subunit chain	
1029	14220702	[Staphylococcus aureus]	1.7
1028	14329/03	[Staphylococcus aureus]	1.7

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 2636683 gb AAC06264,1  (U66333) pol	
1037	2636683	[Schistosoma mansoni]	5.2
l		gi 6016240 sp O02100 HOP1 CAEEL INTEGRAL	
		MEMBRANE PROTEIN HOP-1 pir T15184 presenilin	
	l	beta homolog - Caenorhabditis elegans pir T42237	
		presenilin-beta homolog - Caenorhabditis elegans	
	l .	gb AAB52948,1  (AF000265) Hypothetical protein	
1	· .	C18E3.8 [Caenorhabditis elegans] gb AAB84394.1	
1039	6016240	(AF021905) presenilin [Caenorhabditis elegans]	8.5
		gi 10047173 dbj BAB13380.1  (AB046774) KIAA1554	
1040	10047173	protein [Homo sapiens]	8.4
		gi 14089928 emb CAC13687,1  (AL445564)	
1043	14089928	HEMOLYSIN C [Mycoplasma pulmonis]	4.8
		gi 7469269 pir  S77245 bioY protein - Synechocystis sp.	
		(strain PCC 6803) dbj BAA17579.1  (D90907) BioY	
1045	7469269	protein [Synechocystis sp. PCC 6803]	5.6
		gi 10140780 gb AAG13610.1 AC078840_1	
1046	10140780	(AC078840) hypothetical protein [Oryza sativa]	2.2
	,		
		gi 7496756 pir  T19593 hypothetical protein C31A11.3 -	
		Caenorhabditis elegans emb CAB05685.1  (Z83218)	
		contains similarity to Pfam domain: PF01838 (Domain	
1	1	of unknown function), Score=506.2, E-value=5.4e-154,	
1050	7496756	N=2 [Caenorhabditis elegans]	4.1
		gi 7500420 pir  T32834 hypothetical protein F33H12.1 -	
		Caenorhabditis elegans gb AAB95002.1  (AF040649)	
		Hypothetical protein F33H12.1 [Caenorhabditis	
1052	7500420	elegans]	4
		gi 14736828 ref XP_032481.1  hypothetical protein	
1054	14736828	XP_032481 [Homo sapiens]	5.5
1	700005	gi 7290674 gb AAF46122.1  (AE003436) CG4320 gene	
1056	7290674	product [Drosophila melanogaster]	4.7
1000	1,4050555	gi 14970757 emb CAC44464.1  (AJ313506) CtxX	
1060	14970757	protein [Salmonella typhimurium]	5.5
1000	4115407	gi 4115497 dbj BAA36391.1  (AB010426) AL1 like	2.4
1062	4115497	protein [Phytoplasma sp.]	3.4
1		gi 111816 pir  S21348 probable pol polyprotein-related	
1064	111017	protein 4 - rat emb CAA37647.1  (X53581) ORF4	
1064	111816	[Rattus norvegicus]	6.4

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WC0214500 [fle://E/WO0214500 opc]

	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS			
NO	N	DESCRIPTION	P VALUE	
		gi 14149940 ref[NP_115610.1  hypothetical protein		
1000	1 41 400 40	FLJ23059 [Homo sapiens] dbj BAB15536.1		
1067	14149940	(AK026712) unnamed protein product [Homo sapiens]	8E-34	
		gi 6321761 ref NP_011837,1  Yhl026cp		
		[Saccharomyces cerevisiae] sp P38740 YHC6_YEAST		
		HYPOTHETICAL 30,7 KD PROTEIN IN RIM1-		
		SNF6 INTERGENIC REGION PRECURSOR		
		pir  S48942 hypothetical protein YHL026c - yeast		
1070	C2017C1	(Saccharomyces cerevisiae) gb AAB65062.1  (U11583)		
1072	6321761	YHL026c gene product [Saccharomyces cerevisiae]	0,26	
1076	£022266	gi 5932366 gb AAD56919.1 AF180145_11 (AF180145)		
1076	5932366	hypothetical protein; zm12orf5 [Zymomonas mobilis] gi 7106864 gb AAF36157.1 AF151071 1 (AF151071)	7.2	
1084	7100004		0.0	
1084	7106864		8.3	
		gi 1173846 gb AAA86616.1  (U39455) envelope		
1087	1172046	glycoprotein precursor [Crimean-Congo hemorrhagic fever virus]	6.1	
1087	11/3846		6.1	
		gi 87765 pir  JU0033 hypothetical L1 protein (third		
1092	87765	intron of gene TS) - human prf  1510254A L1 repetitive element ORF [Homo sapiens]	2	
1092	8//03			
		gi 9506813 ref NP_062184.1  Inositol polyphosphate-5- phosphatase [Rattus norvegicus] gb AAB40610.1		
1		(U55192) inositol polyphosphate 5' phosphatase Ship		
1096	9506813	[Rattus norvegicus]	3.3	
1050	2200012	[Kattus noi vogicus]		
		gi 11496734 ref[NP 045511.1  B, burgdorferi predicted		
		coding region BBH18 [Borrelia burgdorferi] pir  C70237		
		hypothetical protein BBH18 - Lyme disease spirochete		
		plasmid H/lp28-3 gb AAC66022.1  (AE000784) B.		
		burgdorferi predicted coding region BBH18 [Borrelia		
1101	11496734	burgdorferi]	7	
1101	*********	gi 9630723 ref NP 047269.1  putative virulence		
		determinant Vir [Mycoplasma arthritidis bacteriophage		
	ĺ	MAV1] gb[AAC33779,1] (AF074945) putative		
		virulence determinant Vir [Mycoplasma arthritidis		
1102	9630723	bacteriophage MAV1]	9.9	
	. 050,25	- oucounclambe v 1		

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS				
NO	N	DESCRIPTION	P VALUE		
		gi 3638957 gb AAC36301.1  (AC004877) sco-spondin-			
		mucin-like; similar to P98167 (PID:g1711548); details			
1107	3638957	of intron/exon structure uncertain [Homo sapiens]	3.3		
		gi 14725582 ref XP_002477.3  hypothetical protein			
1110	14725582	FLJ10829 [Homo sapiens]	0.85		
		gi 7298126 gb AAF53364.1  (AE003642)			
		BG:DS00180.10 gene product [Drosophila			
1112	7298126	melanogaster]	8.6		
		gi 15011489 gb AAK77584.1 AF396436_24			
1120	15011489	(AF396436) heme maturase [Tetrahymena thermophila]	0.077		
		gi 7500306 pir  T21638 hypothetical protein F32B4.1 -			
		Caenorhabditis elegans emb CAB04238.1  (Z81522)			
1121	7500306	predicted using Genefinder [Caenorhabditis elegans]	5.6		
		gi 170156 gb AAA73078.1  (M73688) [Sorghum bicolor			
		endosperm tissue mRNA, complete CDS.], gene product			
1123	170156	prf  1808331A gamma kafirin [Sorghum bicolor]	0.41		
		gi 348951 gb AAC78248.1  (M77194) gag [Rat			
1125	348951	leukemia virus]	4.2		
1					
		gi 14583262 ref NP_127506.1  replicase ORF1ab			
		polyprotein [Equine arteritis virus] emb CAA69187.2			
		(Y07862) replicase ORF1b polyprotein [Cloning vector			
		pEAV030] emb CAC42775.2  (X53459) replicase			
1127	14583262	ORF1b polyprotein [Equine arteritis virus]	0.79		
		gi 6322140 ref NP_012215.1  involved in filamentous			
		growth; Dfg10p [Saccharomyces cerevisiae]			
		sp P40526 YIE9_YEAST HYPOTHETICAL 30.3 KD			
		PROTEIN IN RPL34B-SYG1 INTERGENIC			
		REGION pir S48430 probable membrane protein			
		YIL049w - yeast (Saccharomyces cerevisiae)			
		cmb CAA86173.1  (Z38060) orf, len: 253, CAI; 0.11			
1128	6322140	[Saccharomyces cerevisiae]	7.6		
		gi 14779404 ref XP_008099.4  integrin alpha L			
1129	14779404	precursor [Homo sapiens]	0.004		

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS			
NO	N	DESCRIPTION	P VALUE	
		gi 13928898 rcf NP_113837.1  killer cell lectin-like		
		receptor subfamily G, member 1 [Rattus norvegicus]		
1		pir 159421 mast cell function associated antigen - rat		
		emb CAA56208.1  (X79812) mast cell function		
		associated antigen [Rattus norvegicus]		
		emb CAA65829.1  (X97191) MAFA protein [Rattus		
1136	13928898	norvegicus]	5	
		gi 11467057 ref NP_042533.1  NADH dehydrogenase,		
		subunit 4 [Acanthamoeba castellanii]		
		sp Q37375 NU4M_ACACA NADH-UBIQUINONE		
		OXIDOREDUCTASE CHAIN 4 pir   S53834 NADH		
		dehydrogenase (ubiquinone) (EC 1.6.5.3) chain 4 -		
		Acanthamoeba castellanii mitochondrion		
		gb AAD11826.1  (U12386) NADH dehydrogenase,		
1138	11467057	subunit 4 [Acanthamoeba castellanii]	0.63	
		gi 7331848 gb AAF60536.1  (AC024772) contains		
		similarity to Pfam family PF00569 (Zinc finger present		
		in dystrophin, CBP/p300), score=30.4, E=4.3e-05, N=1		
1140	7331848	[Caenorhabditis elegans]	4.5	
1		gi 12725042 gb AAK06095.1 AE006429_13		
		(AE006429) UNKNOWN PROTEIN [Lactococcus		
1144	12725042	lactis subsp. lactis]	0.82	
		gi 13122173 emb CAC32349.1  (AL583945) putative		
		bifunctional protein (histidine kinase and regulator)		
1146	13122173	[Streptomyces coelicolor]	1.3	
		- 11202555511-114-17740407-114-7251222-1-54-7251222		
	13925661	gi 13925661 gb AAK49407.1 AF261233_1 (AF261233)		
1147	13923661	sodium/calcium exchanger protein [Mus musculus]	5.3	
1149	6552484	gi 6552484 gb AAF16411.1 AF038572_1 (AF038572)	0.02	
1149	0332464	jagged2 [Mus musculus]	0.83	
		gi 13489284 gb AAF16898.2 AF168614_1 (AF168614)		
1154	13489284	HMG-box transcription factor Sox17 [Danio rerio]	9.8	
1101	107204	gil1170115 sp P46430 GTT1 MANSE	7.0	
		GLUTATHIONE S-TRANSFERASE 1 (GST CLASS-		
		THETA) gb AAA92880.1  (L32091) glutathione S-		
1155	1170115	transferase [Manduca sexta]	7.8	
1100		gi 13377412 gb AAK20674.1 AF316639 9 (AF316639)	7.0	
		Wzy [Streptococcus pneumoniae] gb[AAK74527.1]		
		(AE007347) hypothetical protein [Streptococcus		
1159	13377412	pneumoniae]	4,1	
		1		

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
1160	7000060	gi 7298868 gb AAF54075.1  (AE003673) CG1105 gene	
1163	7298868	product [Drosophila melanogaster]	5.5
ļ		gi 10178202 dbj BAB11626.1  (AB016875)	
1164	10178202	gene_id:K9D7.13~unknown protein [Arabidopsis	0.2
1104	10178202	thananaj	9.3
		ail7404370haidC71610 markakla markhana assa biatad	
		gi 7494379 pir  C71610 probable membrane associated protein PFB0615c - malaria parasite (Plasmodium	
		falciparum) gb AAC71912.1  (AE001406) predicted	
1166	7494379	membrane associated protein [Plasmodium falciparum]	6.5
-1100	7.0.075	in a second protein (r. austroaum raiospirum)	
ĺ		gi 12408633 ref NP_074924.1  cytochrome c oxidase	
1		subunit 1 [Podospora anserina]	
1		sp P20681 COX1 PODAN CYTOCHROME C	
		OXIDASE POLYPEPTIDE I pir A48327 cytochrome-c	
ì		oxidase (EC 1.9.3.1) chain I - Podospora anserina	
		mitochondrion emb CAA38777.1  (X55026) cytochrome	
1170	12408633	oxidase c [Podospora anserina]	3.1
		gi 2144233 pir  JC5010 nucleotide-binding protein F -	
		Methanosarcina mazci emb CAA62802.1  (X91502)	
1173	2144233	ABC transporter [Methanosarcina mazei]	6.9
1,174	10501460	gi 10581460 gb AAG20195.1  (AE005096) Vng2034h	0.15
1174	10581460	[Halobacterium sp. NRC-1]	0.15
-		THE STOCKER OF THE COMMON OF T	
		gi 13652647 ref[XP_007000.3  solute carrier family 6 (neurotransmitter transporter, betaine/GABA), member	
ļ		12 [Homo sapiens] ref[XP 029976.1] solute carrier	
İ		family 6 (neurotransmitter transporter, betaine/GABA),	
1176	13652647	member 12 [Homo sapiens]	9
		gi 9631326 ref NP 048159.1  ORF MSV088	
		hypothetical protein [Metanoplus sanguinipes	
l		entomopoxvirus] pir/T28249 ORF MSV088	
	١ .	hypothetical protein - Melanoplus sanguinipes	
	Ι ,	entomopoxvirus gb AAC97639.1  (AF063866) ORF	
		MSV088 hypothetical protein [Melanoplus sanguinipes	
1181	9631326	entomopoxvirus]	3.2
		gi 13897920 gb AAK48502.1 AF260966_1 (AF260966)	
1184	13897920	IL-8 receptor [Oncorhynchus mykiss]	2.6
1,,05	1000000	gi 12860694 dbj BAB32022.1  (AK020187) putative	
1185	12860694	[Mus musculus]	0.86
1107	6671494	gi 6671484 gb AAC49301.2  (U32444) phytochrome F	2.4
1187	00/1484	[Lycopersicon esculentum]	3.4

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundan	Proteins)
SEQ ID	ACCESS		11010
NO	N	DESCRIPTION	P VALUE
		gi 11358531 pir  T51640 myb-related transcription	
		factor MYB19 [imported] - Arabidopsis thaliana	
		(fragment) gb AAC83590.1  (AF062868) putative	
1191	11358531	transcription factor [Arabidopsis thaliana]	0.95
		gi 88462 pir  A27307 proline-rich phosphoprotein (gene	
1192	88462	PRH1, Db allele) - human	7.9
		gi 7484909 pir  T06608 disease resistance protein	
		homolog F16J13.80 - Arabidopsis thaliana	
		cmb CAB40942.1  (AL049638) putative disease	
		resistance protein (TMV N-like) [Arabidopsis thaliana]	
		emb CAB78244.1  (AL161533) putative disease	
1193	7484909	resistance protein (TMV N-like) [Arabidopsis thaliana]	3.4
		gi 7492269 pir  T39663 paired amphipathic helix,	
		probable transcription regulator protein - fission yeast	
		(Schizosaccharomyces pombe) emb CAA21310.1	
1		(AL031856) putative transcriptional rgulatory protein	
1195	7492269	[Schizosaccharomyces pombe]	4.1
		gi 134437 sp P13823 SERA PLAFG SERINE-	
		REPEAT ANTIGEN PROTEIN PRECURSOR (P126)	
		(111 KDA ANTIGEN) pir  A54505 serine-repeat	
'		antigen precursor - malaria parasite (Plasmodium	
		falciparum) (strain FCR3) gb AAA29763.1  (J03993)	
		serine repeat protein [Plasmodium falciparum]	
		gb AAA16791.1  (J04000) serine-repeat antigen protein	
		[Plasmodium falciparum] gblAAA74911.1] (U08113)	
1197	134437	serine repeat antigen [Plasmodium falciparum]	4.6
		gi 12841678 dbj BAB25308.1  (AK007856) putative	
1202	12841678	[Mus musculus]	2E-31
			-
] ]		gi 586120 sp Q07283 TRHY_HUMAN	
		TRICHOHYALIN pir A45973 trichohyalin - human	
1211	586120	gb AAA65582.1  (L09190) trichohyalin [Homo sapiens]	9.8 ،
		gi 3122611 sp O18417 A70A_DROSE ACCESSORY	
		GLAND-SPECIFIC PEPTIDE 70A PRECURSOR	
		(PARAGONIAL PEPTIDE B) cmb CAA67791.1	
1214	3122611	(X99414) sex-peptide [Drosophila sechellia]	6.1
		gi 2190464 emb CAB09537.1  (Z96107) Uncx4.1 [Mus	
1215	2190464	musculus]	7.6

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS			
NO	N	DESCRIPTION	P VALUE	
	i	gi 153930 gb AAA27047.1  (M23007) NADPH-sulfite		
		reductase hemoprotein component [Salmonella		
1216	153930	typhimurium]	1.3	
		gi 5091519 dbj BAA78754.1  (AB023482) Hypothetical		
1217	5091519	protein [Oryza sativa]	0.95	
		gi 9759203 dbj BAB09740.1  (AB015476) heat shock		
	07.0000	transcription factor HSF30-like protein [Arabidopsis		
1219	9759203	thaliana]	9.2	
		gi 1732073 gb AAC50901.1  (U75308) TBP-associated		
1224	1732073	factor [Homo sapiens]	0.13	
		gi 7505421 pir  T23399 hypothetical protein K07C10.1 -		
		Caenorhabditis elegans emb CAA87375.1  (Z47074)		
		similarity to transmembranous domains of the		
		drosophila protein patched (Swiss Prot accession		
1225	7505421	number P18502) [Cacnorhabditis elegans]	3.4	
		gi 140791 sp P17369 YHR3_VACCV	l	
		HYPOTHETICAL HOST RANGE 27.4 KDA		
		PROTEIN pir  WZVZA3 27.4K HindIII-C protein -		
1000	140501	vaccinia virus (strain WR) gb AAA69594.1  (M22812)		
1228	140791	unknown protein [Vaccinia virus]	4,6	
		gi 6831569 sp O84098 IF2_CHLTR TRANSLATION		
		INITIATION FACTOR IF-2 pir H71558 probable		
		translation initiation factor IF-2 - Chlamydia		
1		trachomatis (serotype D, strain UW3/Cx)		
	(001500	gb AAC67687.1  (AE001283) Initiation Factor-2		
1231	6831569	[Chlamydia trachomatis]	7.8	
		gi 12725042 gb AAK06095,1 AE006429_13		
1243	12725042	(AE006429) UNKNOWN PROTEIN [Lactococcus lactis subsp. lactis]	0.01	
1245	12/23042	gil10178036 dbi BAB11519.1  (AB005245)	0,91	
1246	10178036	pectinesterase [Arabidopsis thaliana]	0.26	
1240	10170030		0.28	
		gi 7494298 pir  A71613 hypothetical protein PFB0530c - malaria parasite (Plasmodium falciparum)		
		gb AAC7:1895.1  (AE001400) hypothetical protein		
1249	7494298	[Plasmodium falciparum]	2.6	
1249	1794290	gi 14721018 ref XP 051562.1  similar to agrin (H.	2.0	
1253	14721018	sapiens) [Homo sapiens]	3.5	
1233	1.7/21010	suprous) [1101110 suprous]	3.3	
		  gi 6175163 gb AAF04889.1 AC011437 4 (AC011437)		
1255	6175163	unknown protein [Arabidopsis thaliana]	8.7	
			<u> </u>	

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	D ( ) )
SEQ ID	ACCESS	Table 36 Nearest Neighbor (Blasta vs. Non-Redundant	Proteins)
NO	N N	DESCRIPTION	P VALUE
		gi 12320927 gb AAG50592.1 AC083891 6	T VILLE
	1	(AC083891) ABC transporter, putative [Arabidopsis	
1264	12320927	thalianal	9.5
1201	12,2032,		7.5
	ľ	gi 13375919 ref NP 078940,1  hypothetical protein	
		FLJ11838 [Homo sapiens] dbj BAB13931.1	
1265	12375010	(AK021900) unnamed protein product [Homo sapiens]	4.2
1203	13373919	gi 9369395 gb AAF87143.1 AC002423 8 (AC002423)	4.2
1274	9369395	T23E23.16 [Arabidopsis thaliana]	7.2
12/4	9309393	gi 7435789 pir  T06276 benzothiadiazole-induced	1.2
		protein (clone WCI-4) - wheat gb[AAC49287.1]	
1075	7425700	(U32430) thiol protease [Triticum aestivum]	
1275	7435789	(U32430) tmoi protease [Triticum aestivum]	5.4
		gi 7497192 pir  T19833 hypothetical protein C38D9.3 -	
		Caenorhabditis elegans emb CAB03949.1  (Z81481)	·
1280	7497192	C38D9.3 [Caenorhabditis elegans]	0.15
		gi 14739967 ref XP_035107.1  v-raf murine sarcoma	
1284	14739967	viral oncogene homolog B1 [Homo sapiens]	1.9
		gi 7463772 pir  F70103 signal peptidase I (lepB-1)	
		homolog - Lyme disease spirochete gb AAC66422.1	
		(AE001117) signal peptidase I (lepB-1) [Borrelia	
1289	7463772	burgdorferi]	0.22
	Ì		
		gi 4580388 gb AAD24366.1 AC007171_2 (AC007171)	
1292	4580388	hypothetical protein [Arabidopsis thaliana]	5.5
	1	gi 10437002 dbj BAB14954.1  (AK024666) unnamed	
1293	10437002	protein product [Homo sapiens]	0.000000002
		gi 4493974 emb CAB39033.1  (AL034559) hypothetical	
1296	4493974	protein, PFC0930c [Plasmodium falciparum]	3.2
		gi 482030 pir  S40544 hypothetical protein - Escherichia	
		coli dbj BAA01299.1  (D10483) IS1 hypothetical	
1301	482030	protein E-96(PIR:A04462) [Escherichia coli]	8,3
		gi 4836719 gb AAD30537.1 AF133256_2 (AF133256)	
		envelope protein precursor [Friend mink cell focus-	
1306	4836719	forming virus]	8.6
		gi 12843929 dbj BAB26168.1  (AK009250) putative	
1307	12843929	[Mus musculus]	2.1
		gi 10175999 dbj BAB07095.1  (AP001518)	
1308	10175999	lipopolysaccharide biosynthesis [Bacillus halodurans]	2.8

WO 02/14500

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#### Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins) ACCESS SEO ID NO N DESCRIPTION P VALUE gi|14767733|ref|XP 030859.1| similar to TGF(beta)induced transcription factor 2 (H. sapiens) [Homo 1310 14767733 sapiens] 1.5 gi|1351721|sp|Q10105|YAQ5 SCHPO PUTATIVE TRANSLATIONAL ACTIVATOR C18G6.05C (GCN1 HOMOLOG) pir/T37919 GCN1 homolog fission yeast (Schizosaccharomyces pombe) emb|CAA92385.1| (Z68198) GCN1 homologue 1311 1351721 [Schizosaccharomyces nombe] 0.48 gi|13376747|ref|NP 079428.1| hypothetical protein FLJ12660 [Homo sapiens] ref[XP 017923.1] hypothetical protein FLJ12660 [Homo sapiens] dbj|BAB14203.1| (AK022722) unnamed protein product 1312 13376747 [Homo sapiens] 0.000000003 gi|9453886|dbi|BAB03287.1| (AB045975) pro-alpha 1 1315 9453886 type V/XI collagen [Pagrus major] 0.021 gil11034630|dbi|BAB17154.1| (AP002868) hypothetical protein [Orvza sativa] dbi|BAB55491.1| (AP002541) 1320 11034630 hypothetical protein [Oryza sativa] 0.49 gil14727261|refIXP 027313.1| hypothetical protein FLJ22351 [Homo sapiens] ref[XP 027314.1] 1322 14727261 hypothetical protein FLJ22351 [Homo sapiens] 0.077 gi|2789430|dbj|BAA24380.1| (D30612) repressor 1327 2789430 protein [Homo sapiens] 0.97 gi|7497494|pir||T19963 hypothetical protein C46C2.4 -Caenorhabditis elegans emb|CAA92590.1| (Z68296) contains similarity to Pfam domain; PF00561 (alpha/beta hydrolase fold), Score=30.8, E-value=1e-05. 1328 7497494 N=1 [Caenorhabditis elegans] 0.2 gi|13129018|ref|NP 076956.1| hypothetical protein MGC3077 [Homo sapiens] ref[XP 030116.1] hypothetical protein MGC3077 [Homo saniens] gblAAC23790.11 (AC005154) similar to protein U28928 (PID:g861306) [Homo sapiens] gb|AAH00625.1|AAH00625 (BC000625) Unknown 1329 13129018 (protein for MGC:3077) [Homo sapiens] 2E-89 gi|14647539|gb|AAK71916.1|AF332040 1 (AF332040) 1330 14647539 growth hormone receptor [Jaculus jaculus] 4.8

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteine)
SEQ ID	ACCESS	Tomas Tograd (State 18, 1101 Todatean)	Trotoma)
NO	N	DESCRIPTION	P VALUE
		gi 1584488 prf  2123261AB chemosensory receptor	
1331	1584488	[Caenorhabditis elegans]	0.68
		gi 2326324 emb CAB10936.1  (Z98264) cytochrome c	
1333	2326324	oxidase subunit III [Anabaena sp.]	4.1
	<b>\</b>	gi 7434912 pir H71934 phosphatidylglycerophosphate	
		synthase - Helicobacter pylori (strain J99)	
	ļ	gb AAD05990.1  (AE001475)	
	i	PHOSPHATIDYLGLYCEROPHOSPHATE	
1334	7434912	SYNTHASE [Helicobacter pylori J99]	5.9
	ĺ	gi 1335199 emb CAA26919.1  (X03145) pot. ORF V	
1337	1335199	[Homo sapiens]	5.2
	ļ	gi 7440285 pir  T07994 ribosomal protein S9 -	
		Chlamydomonas reinhardtii chloroplast	
		emb CAA74006.1  (Y13655) 30S ribosomal protein S9	
1340	7440285	[Chlamydomonas reinhardtii]	3
	ĺ	gi 8923088 ref NP_060127.1  hypothetical protein	
		FLJ20080 [Homo sapiens] ref[XP_046636.1]	
		hypothetical protein FLJ20080 [Homo sapiens]	
1041	0000000	dbj BAA90936.1  (AK000087) unnamed protein product	
1341	8923088	[Homo sapiens]	8000000000
		gi 6958206 gb AAF32493.1 AF093132_1 (AF093132)	
1345	6958206	kexin-like protease KEX1 [Pneumocystis carinii f. sp.	2.3
1343	0938200	gi 7496774 pir T32166 hypothetical protein C31B8.8 -	2.3
1346	7496774		2
1340	7420774	Cacionatonis ciogans	
		gi 9366789 emb CAB95551,1  (AL359782) hypothetical	
1350	9366789	protein, CHR1.313. [Trypanosoma brucci]	2.4
	7000.07	gi 10175488 dbi BAB06586,1  (AP001516) cation	
1352	10175488	antiporter (Na+/Ca2+) [Bacillus halodurans]	0.72
		gil1119230 dbi BAA06595,1  (D31786) secretion	
1354	1119230	protein Y [Acyrthosiphon kondoi endosymbiont]	8.5
		gi 2897812 dbi BAA24894.1  (AB010996) G2-G1	
1355	2897812	polyprotein precursor [tomato spotted wilt virus]	8.9
		gi 13111586 gb AAK12388.1 AF296094 1 (AF296094)	
1357	13111586	polyprotein [Porcine teschovirus]	9.2
		gi 13881823 gb AAK46426.1  (AE007064) hypothetical	
1359	13881823	protein [Mycobacterium tuberculosis CDC1551]	1.3
		gi 1362185 pir  S56686 histone H2B123 - wheat	
		dbj BAA07158.1  (D37944) protein H2B123 [Triticum	
1368	1362185	aestivum]	9.7

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	, , , , , , , , , , , , , , , , , , ,	
NO	N	DESCRIPTION	P VALUE
		gi 13812308 ref[NP_113426.1  hypothetical protein	
		[Guillardia theta] emb CAC26995.1  (AJ010592)	
1369	13812308	hypothetical protein [Guillardia theta]	4.7
		gi 13541057 ref[NP_110745.1  Permease (major	
	Ì	facilitator superfamily) [Thermoplasma volcanium]	
		dbj BAB59369.1  (AP000991) unknown product	
1370	13541057	[Thermoplasma volcanium]	7.6
		gi 9964353 rcf NP_064821.1  AMV039 [Amsacta	
		moorei entomopoxvirus] gb AAG02745.1 AF250284_39	
		(AF250284) AMV039 [Amsacta moorei	
1371	9964353	entomopoxvirus]	5.3
i		gi 902377 gb AAA82981.1  (U18059) polyprotein	
1373	902377	[pestivirus type 1]	0.41
·		gi 8953748 dbj BAA98067.1  (AP000368)	
		gene_id:F6B6.1~pir  C71410~similar to unknown	
1374	8953748	protein [Arabidopsis thaliana]	0.27
		·	
		gi 11291752 pir  T47971 seven in absentia-like protein -	
		Arabidopsis thaliana emb CAB71109.1  (AL132959)	
1375	11291752	seven in absentia-like protein [Arabidopsis thaliana]	10
		gi 10177211 dbj BAB10286.1  (AB026650) protein	_
1379	10177211	kinase [Arabidopsis thaliana]	5
		The control of the transport of the control of the	
1204	g0000g0	gi 7332073 gb AAF60760.1  (AC024810) Hypothetical	
1384	7332073	protein Y54E10A.1 [Caenorhabditis elegans]	5.7
		gi 11358814 pir  T46130 RNA polymerase III subunit-	
. 1		like protein - Arabidopsis thaliana emb CAB62010.1  (AL132967) RNA polymerase III subunit-like protein	
1388	11358814	[Arabidopsis thaliana]	2.2
1300	11336614		2.2
		gi 4506569 ref NP_002932.1  roundabout (axon guidance receptor, Drosophila) homolog 1 [Homo	
		sapiens] gb AAC39575.1  (AF040990) roundabout 1	
1389	4506569	[Homo sapiens]	4E-17
1389	4500309	gi 11761072 dbj BAB19062.1  (AP002744) hypothetical	4D-1/
1393	11761072	protein [Oryza sativa]	1.9
1373	11/010/2	Protein for June 1881	1,7
		gi 461649 sp Q05004 BB61 RABIT BRUSH BORDER	
		61.9 KD PROTEIN PRECURSOR pir B45665 adult-	
		specific 61.9K brush border protein precursor - rabbit	
		emb CAA78302.1  (Z12840) protein of unknown	
1398	461649	function [Oryctolagus cuniculus]	8E-52

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESS				
NO	N	DESCRIPTION	P VALUE		
		gi 478809 pir  S29851 protein kinase 6 (EC 2.7.1) -			
		soybean gb AAA34002.1  (M67449) protein kinase			
		[Glycine max] prf  1908223A protein kinase [Glycine			
1402	478809	max]	9.3		
		gi 1517936 gb AAB07000.1  (U52347) tachykinin-like			
1406	1517936	receptor [Stomoxys calcitrans]	1.1		
		gi 9977929 sp Q05013 LIPA_NEIMB CAPSULE			
	!	POLYSACCHARIDE MODIFICATION PROTEIN			
		LIPA pir  D81240 capsule polysaccharide modification			
		protein LipA NMB0082 [imported] - Neisseria			
		meningitidis (group B strain MD58) gb AAF40546.1			
		(AE002367) capsule polysaccharide modification			
1407	9977929	protein LipA [Neisseria meningitidis MC58]	9.3		
1412	14773348	gi 14773348 ref[XP_038450.1  20849 [Homo sapiens]	1E-48		
		gi 13537363 dbj BAB40663.1  (AB051851) death			
1413	13537363	receptor 3 [Homo sapiens]	3.9		
		gi 7477083 pir  A70577 hypothetical protein Rv2133c -			
		Mycobacterium tuberculosis (strain H37RV)			
		emb CAB08660.1  (Z95388) hypothetical protein			
		Rv2133c [Mycobacterium tuberculosis]			
		gb AAK46475.1  (AE007067) conserved hypothetical			
1419	7477083	protein [Mycobacterium tuberculosis CDC1551]	1.9		
		gi 7499021 pir  T20846 hypothetical protein F13E9.9 -			
		Caenorhabditis elegans emb CAA93411.1  (Z69383)			
1422	7499021	F13E9.9 [Caenorhabditis elegans]	4.1		
		gi 14723696 ref XP_035744.1  hypothetical protein			
1424	14723696	XP_035744 [Homo sapiens]	1.5		
		gi 9961349 ref[NP_005500.2  Dmx-like 1 [Homo			
1425	9961349	sapiens]	0.94		
		gi 478302 pir  JN0835 carbonate dehydratase (EC			
1426	478302	4.2.1.1) I - chimpanzee	1.6		
		gi 6649942 gb AAF21641.1 AF032379_1 (AF032379)			
		gonadotrophin releasing hormone receptor; GnRH-R			
1427	6649942	[Trichosurus vulpecula]	9,6		
		gi 6636500 gb AAF20201.1 AF205791_1 (AF205791)			
1432	0036500	squalene synthase [Botryococcus braunii]	1.8		
1424	7504070	gi 7504070 pir  T22586 hypothetical protein F53F4.14 -	0.05		
1434	7504070	Caenorhabditis elegans	0.05		

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WC0214500 [fle://E/WO0214500 opc]__

		Tall an average and average and a	
CTO TD		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	D
NU	IN	DESCRIPTION	P VALUE
		gi 102177 pir  S13141 hypothetical protein (ribosomal	
1437	102177	RNA repeat region) - Giardia lamblia	1.2
	ĺ	gi 13357869 ref NP_078143.1  unique hypothetical	
		[Urcaplasma urealyticum] pir  D82907 hypothetical	
	1	protein UU309 [imported] - Ureaplasma urealyticum	
		gb AAF30718.1 AE002128_6 (AE002128) unique	
1439	13357869	hypothetical [Ureaplasma urealyticum]	5.4
		gi 7468435 pir  B72015 metalloproteinase, insulinase	
	ŀ	family CP0903 [imported] - Chlamydophila pneumoniae	
	ł	(strains CWL029 and AR39) gb AAD19093.1	
		(AE001675) Insulinase family/Protease III	
	[	[Chlamydophila pneumoniae CWL029]	
	1	gb AAF38689.1  (AE002249) mctalloprotease,	
		insulinase family [Chlamydophila pneumoniae AR39]	
		dbj BAA99165.1  (AP002548) insulinase	
1451	7468435	family/protease III [Chlamydophila pneumoniae J138]	7.1
	l	gi 9857712 gb AAG00902.1 AF176776_1 (AF176776)	
		xyloglucan endotransglycosylase LeXET2	
1452	9857712	[Lycopersicon esculentum]	1.9
		gi 4493974 emb CAB39033.1  (AL034559) hypothetical	
1453	4493974	protein, PFC0930c [Plasmodium falciparum]	3.4
	l	gi 4493900 emb CAB39009.1  (AL034558) predicted	
1454		using hexExon; MAL3P2.22 (PFC0265c), Hypothetical	
1454	4493900	protein, len: 637 aa [Plasmodium falciparum]	6.1
		gi 7503603 pir  T16375 hypothetical protein F46G11.1 -	
	ĺ	Caenorhabditis elegans gb AAA81397.1  (U40412)	
		Hypothetical protein F46G11.1 [Caenorhabditis	_
1456	7503603	elegans]	2.1
		gi 9437954 gb AAF87502.1 AF250474_1 (AF250474)	
1460		nucleoprotein [Influenza A virus (A/Duck/Hong	
1462	9437954	Kong/P54/97(H11N9))]	6.7
	1	gi 2129239 pir  G64488 reverse gyrase (intein-	
		containing) - Methanococcus jannaschii	
1464	212222	gb AAB99531.1  (U67592) reverse gyrase, intein	
1464	2129239	containing (rgy) [Methanococcus jannaschii]	9.2
		- 1100000 4001 (BED 000000 111 4 4 4 4 4 4 4 4 4 4 4 4 4 4	
	1	gi 12232439 ref NP_073602.1  hypothetical protein	
1466	12222420	FLJ11937 [Homo sapiens] dbj BAB15124.1	
1466	12232439	(AK025392) unnamed protein product [Homo sapiens]	6.4

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
1468	7243247	gi 7243247 dbj BAA92671.1  (AB037854) KIAA1433 protein [Homo sapiens]	2E-87
1472	12382242	gi 12382242 gb AAG53080.1 AF263824_1 (AF263824) 5'A2rel-related protein [Leishmania donovani]	8.7
1473	13812383	gi 13812383 ref[NP_113501.1  seryl-tRNA synthetase (serin-tRNA ligase) [Guillardia theta] emb CAC27070.1  (AJ010592) seryl-tRNA synthetase (serin-tRNA ligase) [Guillardia theta]	6.5
1476	6014998	gij6014998 spj080164 DPA5_BPR69 DNA POLYMERASE PROCESSIVITY COMPONENT (DNA POLYMERASE ACCESSORY PROTEIN 45) (GP45) pdb IB77 A Chain A, Building A Replisome Structure From Interacting Pieces: A Sliding Clamp Complexed With An Interaction Peptide From Dna Polymerase pdb IB77 B Chain B, Building A Replisome Structure From Interacting Pieces: A Sliding Clamp Complexed With An Interaction Peptide From Dna Polymerase pdb IB77 C Chain C, Building A Replisome Structure From Interacting Pieces: A Sliding Clamp Complexed With An Interaction Peptide From Dna Polymerase pdb IB8H A Chain A, Sliding Clamp, Dna Polymerase pdb IB8H A Chain B, Sliding Clamp, Dna Polymerase pdb IB8H C Alain C, Sliding Clamp, Dna Polymerase gb AAC39310.1  (AP039565) DNA polymerase processivity component [Bacteriophage RB69]	6.9
1170	0014220	gil7706747[ref]NP 057263.1] transient receptor potential	
		gg/ 100/4/picjint_07/205.1 utassient region potential 4 [Homo sapiens] spj@UBN4[TRP4 HUMAN SHORT TRANSIENT RECEPTOR POTENTIAL CHANNEL 4 (TRPC4) (TRP-RELATED PROTEIN 4) (HTRP4) (HTRP4) biJAAD51736, IJAF175406 1	
1479	7706747	(AF175406) transient receptor potential 4 [Homo sapiens] gb AAF22927.1 AF063822_1 (AF063822) trp-related protein 4 [Homo sapiens]	1.7
1480	9757550	gi 9757550 dbj BAB08163.1  (AB030831) SrtT [Streptococcus pyogenes]	0.47
1483	13516917	gi 13516917 dbj BAB40338.1  (AB044076) hybrid sensor [Myxococcus xanthus]	5

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 5790208 dbj BAA83536.1  (AB031285) NADH	
1487	5790208	dehydrogenase subunit 2 [Taenia saginata]	2.7
		gi 14754352 ref XP_032294.1  hypothetical protein	
1489	14754352	FLJ10775 [Homo sapiens]	0.0008
		gi 3552028 gb AAC64946.1  (AF087130) siderophore	
1490	3552028	regulation protein [Neurospora crassa]	7.2
		gi 3913936 sp Q43652 IP27_SOLTU PROTEINASE	
		INHIBITOR TYPE II CM7 PRECURSOR pir   S43105	
		proteinase inhibitor II - potato emb CAA55082.1	
1493	3913936		9.9
		gi 6573777 gb AAF17697.1 AC009243_24 (AC009243)	
1494	6573777	F28K19.17 [Arabidopsis thaliana]	2.2
		gi 13622425 gb AAK34148.1  (AE006569) maltodextrin	
		transport system permease [Streptococcus pyogenes M1	
1496	13622425		3.7
		gi 9910266 ref NP_064627.1  kinesin-like protein 2	
		[Homo sapiens] dbj BAB03309.1  (AB035898) kinesin-	
1498	9910266	like protein 2 [Homo sapiens]	5E-19
		gi 7294128 gb AAF49482.1  (AE003527) CG4925 gene	
1503	7294128	product [Drosophila melanogaster]	3.6
		gi 14089610 emb CAC13370.1  (AL445563) unknown;	
1506	14000010	predicted coding region [Mycoplasma pulmonis]	1.5
1306	14089610	gi 2565196 gb AAB81938.1  (AF000381) non-functional	1.3
1507	2565196	folate binding protein [Homo sapiens]	0.0000004
1307	2303190	gi 14753935 ref XP 040892.1  hypothetical protein	0.000004
1509	14753935		3.3
1509	14733933		3,3
		gi 12545425 ref NP_074975.1  hypothetical protein	
		[Astasia longa] sp[P34776]YCY2_ASTLO	
	1	HYPOTHETICAL 34.5 KDA PROTEIN IN RPS12- TRNP INTERGENIC REGION (ORF288)	
	i	emb CAC24586.1  (AJ294725) hypothetical protein	
1510	12545425		7
1310	12343423	[Asiasia ionga]	
1511	420215	gi 420215 pir  B45878 hypothetical protein 2 - mouse	7.1
	.202.0	gi 7512874 pir  T08792 hypothetical protein	
		DKFZp586E1422.1 - human (fragment)	
		emb CAB43306.1  (AL050170) hypothetical protein	
1512	7512874	[Homo sapiens]	5.5
	1	gi 221758 dbi BAA01683.1  (D10879) UL37 [human	
1514	221758	herpesvirus 1]	0.18

	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS			
NO	N_	DESCRIPTION	P VALUE	
ļ				
		gi 11466232 ref[NP_062855.1  ORF9, contains 8 trans		
		membrane regions, putative [Physarum polycephalum]		
ļ.		dbj BAB08089.1  (AB027295) ORF9, contains 8 trans		
1517	11466232	membrane regions, putative [Physarum polycephalum]	0.43	
		gi 182710 gb AAA52467,1  (M13918) fibronectin		
1518	182710	receptor alpha-subunit precursor [Homo sapiens]	1.3	
		gi 1142976 gb AAC52392.1  (U28769) odorant receptor		
		[Mus musculus] prf  2207403C odorant receptor [Mus		
1519	1142976	musculus]	5.6	
	l			
		gi 6912446 ref NP_036417.1  potassium voltage-gated		
		channel, subfamily H (eag-related), member 4; ether-a-		
		go-go K(+) channel family member [Homo sapiens]		
1520	6912446	dbj BAA83592.1  (AB022698) BEC2 [Homo sapiens]	8.4	
		gi]14601483 ref[NP_148021.1  hypothetical protein		
		[Aeropyrum pernix] pir  G72637 hypothetical protein		
1	1	APE1558 - Aeropyrum pernix (strain K1)		
		dbj BAA80557.1  (AP000061) 279aa long hypothetical		
1522	14601483	protein [Aeropyrum pernix]	5.7	
ì		Bearing III I Great III Too and A Caro		
		gi 12515306 gb AAG56369.1 AE005365_3 (AE005365)		
		orf, hypothetical protein [Escherichia coli O157:H7 EDL933] dbijBAB35435.1] (AP002557) hypothetical		
1526	12515206	protein [Escherichia coli O157:H7]	7.4	
1320	12313300	gi 2765672 cmb CAB06819.1  (Z86115) ArbX	7.4	
1530	2765672	[Lactobacillus delbrueckii]	7.9	
1330	2703072	gi 13472514 ref NP 104081.1  unknown protein	1.5	
		[Mesorhizobium loti] dbi BAB49867.1  (AP003000)		
1532	13472514	unknown protein [Mesorhizobium loti]	7.9	
1332	13472314	gi 3309522 gb AAC26098.1  (U18292) unknown	1.2	
1533	3309522	[Borrelia burgdorferi]	2.1	
1555	55 65 62 2	[2010th Dangarien]		
		gi 14520328 ref[NP 125803.1  activator 1, replication		
	Į.	factor C, small subunit [Pyrococcus abyssi] pir  C75198		
	Į	activator 1, replication factor c, small chain PAB0068 -		
		Pyrococcus abyssi (strain Orsay) emb CAB49034.1		
	1	(AJ248283) activator 1, replication factor C, small		
1535	14520328	subunit [Pyrococcus abyssi]	0.41	
	1	gi/7229605 gb AAF42902.1  (AF229961) NADH		
1536	7229605	dehydrogenase subunit 1 [Taygetis andromede]	6.7	

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 6681261 ref[NP_031926.1  ect2 oncogene [Mus	
		musculus] sp Q07139 ECT2_MOUSE ECT2 PROTEIN	
		(ECT2 ONCOGENE) pir  S32372 transforming protein	
		(ect2) - mouse gb AAA37536.1  (L11316) ect2 [Mus	
		musculus] prf  1911407A oncogene cct2 [Mus	
1537	6681261	musculus]	6E-10
		gi 12381848 emb CAC24715.1  (AJ297319) glucose-6-	
		phosphate dehydrogenase-6-phosphogluconolactonase	
1538	12381848	[Plasmodium berghei]	7
		gi 7688657 gb AAF67469.1 AF146760_1 (AF146760)	
1540	7688657	septin 2-like cell division control protein [Homo sapiens]	9.7
		gi 13021853 gb AAK11564.1 AF318500_1 (AF318500)	
1541	13021853	ent-kaurenoic acid hydroxylase [Arabidopsis thaliana]	0.091
		gi 6900006 emb CAB71294.1  (AJ251917) chorion	
1543	6900006	protein s18 [Ceratitis capitata]	. 2.1
		gi 1076445 pir  S53004 mitosis-specific cyclin CYC2 -	
1545	1076445	rape gb AAA51660.1  (L25406) cyclin [Brassica napus]	4.3
1	1	gi 7293274 gb AAF48655.1  (AE003503) CG9644 gene	
1548	7293274	product [Drosophila melanogaster]	0.46
1		gi 11358961 pir  T51243 Scl1 protein [imported] - rice	
		(fragment) gb AAC98091.1  (AF067401) Scl1 protein	
1549	11358961	[Oryza sativa]	3.9
		gi 8978966 dbj BAA98801.1  (AP002547) phenylalanyl	
		tRNA synthetase beta [Chlamydophila pneumoniae	
1550	8978966	J138]	4.1
		gi 14010341 gb AAK51958.1 AF362013_1 (AF362013)	
		ATP synthase F0 subunit 6 [Halichondria sp. RFW-	
1551	14010341	2001]	8.2
		gi 8923094 ref NP_060130.1  hypothetical protein	
		FLJ20085 [Homo sapiens] ref[XP_009383,2]	
		hypothetical protein FLJ20085 [Homo sapiens]	
		dbj BAA90939.1  (AK000092) unnamed protein product	
1553	8923094	[Homo sapiens]	1.6
		gi 14724725 ref XP_037600.1  hypothetical protein	
1557	14724725	XP_037600 [Homo sapiens]	4.2

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS N	DUG (D. 1997)	
NO	l N	DESCRIPTION	P VALUE
		gi 14758684 ref XP_007211.4  retinoblastoma 1	
		(including osteosarcoma) [Homo sapiens]	
		ref[XP_033825.1] retinoblastoma 1 (including	
		osteosarcoma) [Homo sapiens] ref[XP_033826.1]	
		retinoblastoma 1 (including osteosarcoma) [Homo	
1561	14758684	sapiens] ref XP_033827.1  retinoblastoma 1 (including osteosarcoma) [Homo sapiens]	0.91
1301	14730064	osteosarcoma) [riomo sapiens]	0.91
		gi 9635387 ref NP 059285.1  ORF137 [Xestia c-nigrum	
		granulovirus] gb AAF05251,1 AF162221 137	
1565	9635387	(AF162221) ORF137 [Xestia c-nigrum granulovirus]	5,9
1505	7055507	(12 102221) Old 137 (Mediat Congram granatovitus)	3.9
		gi 1730077 sp P18160 KYK1 DICDI NON-	
		RECEPTOR TYROSINE KINASE SPORE LYSIS A	
		(TYROSINE-PROTEIN KINASE 1) pir T18276 non-	
		receptor tyrosine kinase - slime mold (Dictyostelium	
		discoideum) gb AAB41125,1  (U32174) non-receptor	
1572	1730077	tyrosine kinase [Dictyostelium discoideum]	6
		gi 9622133 gb AAF89633.1 AF167719 1 (AF167719)	
1574	9622133	transmembrane leptin receptor [Sus scrofa]	2.6
		gi 139809 sp P27571 XIST MOUSE X INACTIVE	
		SPECIFIC TRANSCRIPT PROTEIN pir  S15433	
		hypothetical protein - mouse emb CAA41978.1	
		(X59289) ORF [Mus musculus] prf  1711440A xist	
1578	139809	gene [Mus musculus]	7.7
		gi 8567792 gb AAF76364.1  (AC013428) I-box binding	
1579	8567792	factor, putative [Arabidopsis thaliana]	5.7
		gi 7638161 gb AAF65408.1 AF238312_1 (AF238312)	
		putative serine-threonine protein kinase MkcB	
1581	7638161	[Dictyostelium discoideum]	9.2
		Nacities loogedelayer can be	
		gi 12644495 sp Q9Z7G7 EX5B_CHLPN	
		EXODEOXYRIBONUCLEASE V BETA CHAIN	
1583	12644405	dbj BAA98945.1  (AP002547) exodeoxyribonuclease V,	7.2
1363	12044493	beta [Chlamydophila pneumoniae J138]	7.2
		gi 11350667 pir  D83160 nitrite extrusion protein 1	
		PA3877 [imported] - Pseudomonas aeruginosa (strain	
		PAO1) emb CAA75538.1  (Y15252) nitrate extrusion protein [Pseudomonas acruginosa]	
		gb AAG07264.1 AE004805 2 (AE004805) nitrite	
1588	11350667	extrusion protein 1 [Pseudomonas aeruginosa]	9.7
1500	11330007	extrusion protein 1 [r settuomonas acruginosa]	9.1

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESS				
NO	N	DESCRIPTION	P VALUE		
		gi 12644495 sp Q9Z7G7 EX5B_CHLPN			
		EXODEOXYRIBONUCLEASE V BETA CHAIN			
1500		dbj BAA98945.1  (AP002547) exodeoxyribonuclease V,			
1589	12644495	beta [Chlamydophila pneumoniae J138]	5,2		
		  gi 305479 gb AAC37807.1  (L11473) envelope			
1590	305479	glycoprotein [Human immunodeficiency virus type 1]	4.3		
1390	303479	grycoprotein (riuman minimunouenciency virus type 1)	4.3		
		gi 9055244 ref NP 061256.1  huntington yeast partner C			
1	ł	[Mus musculus] gb AAD39464.1 AF135440 1			
1591	9055244	(AF135440) huntington yeast partner C [Mus musculus]	0.8		
	7000211	gi 12666210 emb CAC28083.1  (AL138875)	V.0		
1593	12666210	bA103J18.2 (novel protein) [Homo sapiens]	4		
		gi 12855510 dbi BAB30362.1  (AK016654) putative			
1596	12855510	[Mus musculus]	0.067		
		gi 133747 sp P09899 RS12 MICLU 30S RIBOSOMAL			
		PROTEIN S12 pir A26956 ribosomal protein S12 -			
		Micrococcus luteus gb AAA25317.1  (M17788)			
		ribosomal protein S12 (gtg start codon) [Micrococcus			
1597	133747	luteus]	9.9		
		gi 7292455 gb AAF47859.1  (AE003480) CG15005			
1600	7292455	gene product [Drosophila melanogaster]	0.43		
		gi 14748674 ref XP_038133.1  Prader-Willi/Angelman			
1601	14748674	syndrome-5 [Homo sapiens]	8.9		
		gi 11278020 pir  H82215 serine transporter VC1301			
		[imported] - Vibrio cholerae (group O1 strain N16961)			
1602	11278020	gb AAF94460.1  (AE004210) serine transporter [Vibrio			
1002	11278020	gi 9757538 dbi BAB08122.1  (AB030852) maturase	7.8		
1607	9757538		6.6		
1507	2131336	gi 12249143 ref NP 066194.2  NADH dehydrogenase	0.0		
		subunit 5 [Schistosoma japonicum] gblAAG13134,2			
		(AF215860) NADH dehydrogenase subunit 5			
1618	12249143	[Schistosoma japonicum]	5.9		
		The second secon			
]		gi 7495508 pir  T18993 hypothetical protein C06B8,1 -			
		Caenorhabditis elegans emb[CAB03850,1] (Z81463)			
		Similarity to C.elegans zinc finger proteins, contains			
		similarity to Pfam domain: PF00104 (Ligand-binding			
		domain of nuclear hormone receptor), Score=-13.7, E-			
1620	7495508	value=0.051, N=1 [Caenorhabditis elegans]	2.9		

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
l		gi 7288570 gb AAF45170.1 AF218963_1 (AF218963)	
1621	7288570	period [Drosophila miranda]	4.3
		gi 7707666 dbj BAA95343.1  (AB027560) ATPase	
1622	7707666	subunit 6 [Echinococcus vogeli]	4.1
	ĺ		
		gi 4761646 gb AAD29428.1 AF139060_1 (AF139060)	
	l	transmembrane cell adhesion receptor MUA-3 precursor	
		[Caenorhabditis elegans] emb CAA83226.2  (Z30974)	
		contains similarity to Pfam domain: PF00057 (Low-	
		density lipoprotein receptor domain class A),	
	l	Score=52.9, E-value=2.2e-12, N=3, second half (former	
		T20G5.3) contains similarity to Pfam domain: PF00008	
		(EGF-like domain), Score=326.3, E-value=1>	
		emb CAC42345.1  (Z30423) contains similarity to Pfam domain: PF00057 (Low-density lipoprotein receptor	
		domain: Proous / (Low-density lipoprotein receptor domain class A), Score=52.9, E-value=2.2e-12, N=3,	
		second half (former T20G5.3) contains similarity to	
1		Pfam domain: PF00008 (EGF-like domain),	
1625	4761646	Score=326.3, E-value=1>	1.7
1		gi 3582424 dbj BAA33057.1  (AB017255) arginine	
1626	3582424	kinase two-domain chain [Pseudocardium sachalinensis]	4.5
		gi 9758077 dbj BAB08521.1  (AB009052) sucrose	
1628	9758077	cleavage protein-like [Arabidopsis thaliana]	6.2
		gi 11610630 gb AAG37436.1  (AY013711) cyclin E	
1629	11610630	[Mustela vison]	10
		gi 4838093 gb AAD30838.1  (AF103278)	
		immunoglobulin heavy chain variable region [Homo	
1631	4838093	sapiens]	3,3
1		gi 12045265 ref[NP_073076.1  ATP synthase F0,	
		subunit B (atpF) [Mycoplasma genitalium]	
		sp P47643 ATPF_MYCGE ATP SYNTHASE B	
		CHAIN PRECURSOR pir  F64244 ATP synthase B	
		chain (atpF) - Mycoplasma genitalium gb AAC71631.1	
1,000	100 450	(U39722) ATP synthase F0, subunit B (atpF)	
1635	12045265	[Mycoplasma genitalium]	8.5
1636	627406	gi 627406 pir  A54849 collagen alpha 1(VII) chain	0.66
1030	627406	precursor - human	0.66
1643	2246540	gi 2246540 gb AAB62665.1  (U93872) glycoprotein M [Human herpesvirus 8]	1.5
1043	440340	gi 2388576 gb AAB71457.1  (AC000098) YUP8H12.17	1.3
1644	2388576	[Arabidopsis thaliana]	4.7
	2200270	[Lataordopais manama]	7.7

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 7490184 pir  T37997 carboxypeptidase y - fission	
		yeast (Schizosaccharomyces pombe) pir  T43236	
		carboxypeptidase C (EC 3.4.16.5) precursor [validated]	
		fission yeast (Schizosaccharomyces pombe)	
		emb CAB10121.1  (Z97209) carboxypeptidase y	
		[Schizosaccharomyces pombe] dbj BAA25568.1	
		(D86560) carboxypeptidase Y [Schizosaccharomyces	
1645	7490184		0.009
		gi 7415597 dbj BAA93452.1  (AB026494)	
1646	7415597	acyltransferase homolog [Gentiana triflora]	6.1
		gi 13541404 ref[NP_111092.1  DNA helicase	
		(superfamily II) [Thermoplasma volcanium]	
		dbj BAB59714.1  (AP000992) DNA helicase	
1648	13541404	[Thermoplasma volcanium]	7.5
		gi 6174902 sp Q24767 PER_DROYA PERIOD	
		CIRCADIAN PROTEIN pir  S17286 period clock	
		protein - fruit fly (Drosophila yakuba)	
		emb CAA43439.1  (X61127) period [Drosophila	
1649	6174902	yakuba]	0.2
		gi 14318508 ref NP_116641.1  Ies1p [Saccharomyces	
		cerevisiae] sp P43579 YFB3_YEAST	
		HYPOTHETICAL 78.8 KD PROTEIN IN HSP12-	
		HXT10 INTERGENIC REGION pir  S48316 probable	
		membrane protein YFL013c - yeast (Saccharomyces	
		cerevisiae) emb CAA86347.1  (Z46255) orf, len: 692,	
		CAI: 0.14 [Saccharomyces cerevisiae] dbj BAA09225.1	
1650	14318508	(D50617) YFL013C [Saccharomyces cerevisiae]	8.8
		gi 5052950 gb AAD38784.1 AF149422_1 (AF149422)	
1655	5052950	unknown [Homo sapiens]	0.000000007
		gi 2988422 gb AAC39776.1  (AF016903) agrin	
1660	2988422	precursor [Homo sapiens]	3.4
1661	14756100	gi 14756108 ref XP_029883.1  EGF-like-domain,	_
1661	14/56108	multiple 4 [Homo sapiens]	5
1004	10047212	gi 10047313 dbj BAB13444.1  (AB046838) KIAA1618	0.00001
1664	1004/313	protein [Homo sapiens]	0.00001
		Usersand Chill consent strawers	
		gi 5902048 ref NP_008974.1  HIV-1 rev binding protein	
		2; Rev interacting protein [Homo sapiens] pir  G02629	
1669	5003040	Rev interacting protein Rip-1 - human gb AAB00557.1	0.000
1009	3902048	(U55766) Rev interacting protein Rip-1 [Homo sapiens]	0.009

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SEQ ID   ACCESS NO   DESCRIPTION   P VALUE			Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
1672   13122465   g 13122465   g 705cin-coupled receptor   Flomo sapiens    2.4	SEQ ID	ACCESS		
1672   13122465   G protein-coupled receptor [Homo sapiens]   2.4	NO	N	DESCRIPTION	P VALUE
1672   13122465   G protein-coupled receptor [Homo sapiens]   2.4	1			
gi 7296752 gb AAF52030.1  (AE003603) CG12147   2.1     1673   7296752  gene product [Drosophila melanogaster]   2.1     1681   14767950 reffXP_040494.1  nuclear RNA export factor 5 [Homo sapiens]   6.9     1681   14767950  nuclear RNA export factor 5 [Homo sapiens]   6.9     1681   14767950  nuclear RNA export factor 5 [Homo sapiens]   6.9     1682   14740021 reffXP_033955.1  32575 [Homo sapiens]   7     1684   14740021 reffXP_033955.1  32575 [Homo sapiens]   7     1685   13568988   gbhAAK30843.1 AF254571_1 (AF254571)   7     1685   13568988   growth/differentiation factor 7 [Mus musculus]   5.8     1694   106323   piir A34087 hypothetical protein (L1H 5' region) - human   9     106323   region) - human   9     106324   region) - human   9     106325   region) - human   9     106326   region) - human   9     106327   region) - human   9     106328   region) - human   9     106329   reffXP_012087 gag/pol polyprotein maize retrotransposon Hopscotch gb AAA57005.1  (U12626)   0     1065   7444442   piir T02087 gag/pol polyprotein [Zea   0     1076   13569915   reffXP_112205.1  amuionless protein   (Homo sapiens)   0     1076   13569915   (AF238788) amuinokses [Homo sapiens]   0     1077   13812078   reffXP_113215.1   hypothetical protein   (Diullardia theta)   0     1077   13812078   reffXP_113215.1   hypothetical protein   (Diullardia theta)   0     1078   reffXP_079359.1   hypothetical protein   0     1079   reffXP_079359.1   hypothetical protein   0     1070   13376638   reffXP_079359.1   hypothetical protein   0     170   13376638   (AK024781) unnamed protein product [Homo sapiens]   5     170   13376638   (AK024781) unnamed protein product [Homo sapiens]   5     170   17445803   difficile   0   0   28     1813953331   dbj BAA34722.1   (AB015856) ATF6   0   0   0   0     1075016   BH2396-unknown [Bacillus halodurans]   9     10175016   BH2396-unknown [Bacillus halodurans]   9     10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016	1670	12122465		
1673   7296752   gene product [Drosophila melanogaster]   2.1	1672	13122403		2.4
gi 14767950 ref XP_040494.1  unclear RNA export factor 5 [Homo sapiens] emb CAC20428.1  (AJ277654)   muclear RNA export factor 5 [Homo sapiens]   6.9   gi 1474002 ref XP_033953.1  32575 [Homo sapiens]   ref XP_033954.1  32575 [Homo sapiens]   2.5   ref XP_033954.1  32575 [Homo sapiens]   2.5   ref XP_033954.1  32575 [Homo sapiens]   2.5   ref XP_033954.1  32575 [Homo sapiens]   2.5   ref XP_033955.1  32577 [Homo sapiens]   3.568988   gowth/differentiation factor 7 [Mus musculus]   5.8   ref XP_033955.1  32577 [Homo sapiens]   3.68988   rowth/differentiation factor 7 [Mus musculus]   5.8   ref XP_0323  region) - human   0.005   ref XP_0323  region) - human   0.005   ref XP_0323  region) - human   0.005   ref XP_0444442  ref XP_03231  ref XP_0444442  ref XP_03231  ref XP_0444442  ref XP_03231  ref XP_0444442  ref XP_03231  ref XP_0444442  ref XP_03231  ref XP_0444442  ref XP_03434  ref XP_04444442  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_0444444  ref XP_04444444  ref XP_04444444  ref XP_04444444  ref XP_04444444  ref XP_04444444  ref XP_04444444  ref XP_04444444  ref XP_04444444  ref XP_04444444  ref XP_04444444  ref XP_04444444  ref XP_04444444  ref XP_04444444  ref XP_04444444  ref XP_04444444  ref XP_044444444  ref XP_04444444  ref XP_04444444  ref XP_044444444  ref XP_044444444  ref XP_04444444  ref XP_044444444  ref XP_044444444  ref XP_04444444  r	1673	7206752		2.1
factor 5 [Homo sapiens] emb[CAC20428.1] (AJ277654)     14767950   nuclear RNA export factor 5 [Homo sapiens]   6.9     spil14740021 [refRP_033954.1] 32576 [Homo sapiens]     refRP_033954.1] 32576 [Homo sapiens]     refRP_033954.1] 32576 [Homo sapiens]     refRP_033954.1] 32576 [Homo sapiens]     1684   14740021 refRP_033955.1] 32577 [Homo sapiens]     refRP_033955.1] 32577 [Homo sapiens]     1685   13568988 gbhAK30843.1]AF254571_1 (AF254571)     1686   13568988 gbhAK30843.1]AF254571_1 (AF254571)     1694   106323 region) - human     spil1444442 [pir][T02087 gag/pol polyprotein - maize retrotransposon Hopscotch gbhAAA57005.1] (U112626)     copia-like retrotransposon Hopscotch polyprotein [Zea mays]     0.49     gil13569915 [refRP_112205.1] amnionless protein     Homo sapiens] gbhAK28352.1]AF328788_1     1696   13569915 [AF328788] amnionless [Homo sapiens]   4.4     gil13569915 [AF328788] amnionless [Homo sapiens]     1697   13812078 [AF3083031) hypothetical protein [Guillardia theta]     1698   2462193 [spiQ06603 [GOLI_DROME GOLIATH PROTEIN (GI PROTEIN) pir][C1495 regulatory protein GI - fruit fly (Droscophila melanogaster)     gil13376638 [refRP_079359.1] hypothetical protein [Drosophila melanogaster]     1702   13376638 [AK024781] unamend protein protein [TDrosophila melanogaster]     1703   7445803 [pir][C348 cdd4 protein - Clostridium difficile emb[CAA63569.1] (X92982) cdd4 [Clostridium difficile]     1705   7445803 [difficile]     1706   10175016 [BH2396-unknown [Bacillus halodurans]     gil3953531 [dbjBAA343722.1] (AB015856) ATF6	1073	1290132	gene product [Drosophila metanogaster]	Z,1
factor 5 [Homo sapiens] emb[CAC20428.1] (AJ277654)     14767950   nuclear RNA export factor 5 [Homo sapiens]   6.9     spil14740021 [refRP_033954.1] 32576 [Homo sapiens]     refRP_033954.1] 32576 [Homo sapiens]     refRP_033954.1] 32576 [Homo sapiens]     refRP_033954.1] 32576 [Homo sapiens]     1684   14740021 refRP_033955.1] 32577 [Homo sapiens]     refRP_033955.1] 32577 [Homo sapiens]     1685   13568988 gbhAK30843.1]AF254571_1 (AF254571)     1686   13568988 gbhAK30843.1]AF254571_1 (AF254571)     1694   106323 region) - human     spil1444442 [pir][T02087 gag/pol polyprotein - maize retrotransposon Hopscotch gbhAAA57005.1] (U112626)     copia-like retrotransposon Hopscotch polyprotein [Zea mays]     0.49     gil13569915 [refRP_112205.1] amnionless protein     Homo sapiens] gbhAK28352.1]AF328788_1     1696   13569915 [AF328788] amnionless [Homo sapiens]   4.4     gil13569915 [AF328788] amnionless [Homo sapiens]     1697   13812078 [AF3083031) hypothetical protein [Guillardia theta]     1698   2462193 [spiQ06603 [GOLI_DROME GOLIATH PROTEIN (GI PROTEIN) pir][C1495 regulatory protein GI - fruit fly (Droscophila melanogaster)     gil13376638 [refRP_079359.1] hypothetical protein [Drosophila melanogaster]     1702   13376638 [AK024781] unamend protein protein [TDrosophila melanogaster]     1703   7445803 [pir][C348 cdd4 protein - Clostridium difficile emb[CAA63569.1] (X92982) cdd4 [Clostridium difficile]     1705   7445803 [difficile]     1706   10175016 [BH2396-unknown [Bacillus halodurans]     gil3953531 [dbjBAA343722.1] (AB015856) ATF6			gil 14767950 reffXP 040494 1  nuclear RNA export	
14767950   muclear RNA export factor 5 [Homo sapiens]   6.9   gi]14740021   refkP. 033954.   132576 [Homo sapiens]   refkP. 033954.   132576 [Homo sapiens]   2.5   refkP. 033954.   132576 [Homo sapiens]   2.5   refkP. 033954.   132576 [Homo sapiens]   2.5   gi]1356898.   gi]1356898.   gi]1356898.   gi]1356898.   gi]1356898.   gi]1356898.   gi]1356898.   gi]1356898.   gi]1356898.   gi]1356898.   gi]1356898.   gi]1356898.   gi]1356898.   gi]135498.   gi]135498.   gi]135498.   gi]135498.   gi]135498.   gi]135498.   gi]135498.   gi]135498.   gi]135498.   gi]135498.   gi]135498.   gi]1356991.   gi]1356991.   gi]1356991.   gi]1356991.   gi]1356991.   gi]1356991.   gi]1356991.   gi]1358991.   gi]1358991.   gi]1358991.   gi]1358991.   gi]1358991.   gi]1358991.   gi]1358991.   gi]1358991.   gi]1358991.   gi]1358991.   gi]135891.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.   gi]1381.				
gi 14740021 refiXP_033953.1  32575 [Homo sapiens] refiXP_033954.1  32576 [Homo sapiens] refiXP_033954.1  32576 [Homo sapiens]   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.5   2.	1681	14767950		6.9
reflXP, 033954, 1] 32576 [Homo sapiens]   2.5     1684   14740021   reflXP, 033955.1] 32577 [Homo sapiens]   2.5     1685   1356898 growth/differentiation factor 7 [Mus musculus]   5.8     1686   106323   region) - human   5.8     1694   106323   region) - human   0.005     1695   reflored   166323   region) - human   0.005     1696   106323   region) - human   0.005     1697   reflored   166323   region) - human   0.005     1698   reflored   166323   region) - human   0.005     1699   reflored   166323   reflored   1695   reflored   1695   reflored   1695   reflored   1695   reflored   1695   reflored   1695   reflored   1695   reflored   1696   13569915   reflored   1205.1]   amnioaless protein   1696   13569915   reflored   13569915   reflored   1352.1]   reflored   1696   13569915   reflored   1352.1]   reflored   1696   13812078   reflored   13812078   reflored   13812078   reflored   13812071   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   1697   reflored   169				
gi 13568988 gb AAK30843.1 AF254571_1 (AF254571)     1685   13568988  growth/differentiation factor 7 [Mus musculus]   5.8     1694   106323  gi 106323 pir  A34087 hypothetical protein (L1H 5' region) - human   0.005     1695   gi 7444442 pir [T02087 gag/pol polyprotein - maize retrotransposon Hopscotch gb AAA57005.1  (U12626)     1696   copia-like retrotransposon Hopscotch polyprotein [Zea mays]   0.49     1697   13669915 refNP_112205.1  amuionless protein   Homo sapiens  gb AAK28332.1 AF328788_1     1698   13569915  (AF328788) amuionless [Homo sapiens]   4.4     1699   1356915  (AF328788) amuionless [Homo sapiens]   4.4     1690   1356915  (AF328788) amuionless [Homo sapiens]   5.2     1691   13812078  (AF083031) hypothetical protein [Guillardia theta]   5.2     1692   13812078  (AF083031) hypothetical protein [Guillardia theta]   5.2     1693   13812078  (AF083031) hypothetical protein [Guillardia theta]   5.2     1694   13812078  (AF083031) hypothetical protein [Drosophila melanogaster]   9.3     1699   462193   melanogaster]   9.3     1699   462193   melanogaster]   9.3     1700   13376638   (AK024781) unnamed protein protein [Drosophila melanogaster]   9.3     1701   13376638   (AK024781) unnamed protein protein [Homo sapiens]   5E-49     1702   13376638   (AK024781) unnamed protein protein [Homo sapiens]   5E-49     1705   7445803   difficile   0.28   (1792982) cdd4 [Clostridium difficile emb CAA63569.1  (X92982) cdd4 [Clostridium difficile]   0.28   (179204)   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10175016   10				
1685   13568988 growth/differentiation factor 7 [Mus musculus]   5.8	1684	14740021	ref[XP_033955.1] 32577 [Homo sapiens]	2.5
1685   13568988 growth/differentiation factor 7 [Mus musculus]   5.8				
1694   106323				
1694   106323   region) - human   0.005	1685	13568988		5.8
gji7444442 pir  T02087 gag/pol polyprotein - maize retrotransposon Hopscotch gb AAA57005.1 (U12626)				
retrotransposon Hopscotch gb AAA57005.1  (U12626)   copia-like retrotransposon Hopscotch polyprotein [Zea copia-like retrotransposon Hopscotch polyprotein [Zea copia-like retrotransposon Hopscotch polyprotein [Zea copia-like retrotransposon Hopscotch polyprotein [Zea gil13569915]   classification   classifica	1694	106323		0.005
1695   7444442				
1695   7444442   mays    0.49				
gij13569915 ref NP_112205.1  amnionless protein	1605	744444		0.40
Iffomo sapiens  gb AAK28532.1 AF528788_1     13569915 (AF328788) ammionless [Homo sapiens]   4.4     13569915 (AF328788) ammionless [Homo sapiens]   4.4     13812078 (AF328788) ammionless [Homo sapiens]     1697   13812078 (AF083031) hypothetical protein [Guillardia theta]   5.2     13812078 (AF083031) hypothetical protein [Guillardia theta]   5.2     13812078 (AF083031) hypothetical protein [Guillardia theta]   7.2     13812078 (AF083031) hypothetical protein [Guillardia theta]   7.2     13812078 (AF08382.1] (M97204) goliath protein [Drosophila melanogaster)   9.3     1699   462193   1376638   14097204) goliath protein [Drosophila melanogaster]   9.3     1702   13376638   1376638   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204   14097204	1093	744442		0.49
13569915 (AF328788) amnionless [Homo sapiens]   4.4				
gi  3812078 ref NP_113215.1  hypothetical protein   Guillardia thetal gb AAF24011.1 AF083031_8   S45830331) hypothetical protein [Guillardia thetal gb AAF24011.1 AF083031_8   S45830331) hypothetical protein [Guillardia thetal gb AAF24011.1 AF083031_8   S4583031] hypothetical protein [Guillardia thetal gb AAF24013 sp Q06003 GOLI_DROME GOLIATH   PROTEIN (GI PROTEIN) pir  JI C1495 regulatory   protein GI - fruit fly (Droscophia melanegaster)   gb AAA28582.1  (M97204) goliath protein [Drosophila melanogaster]   9.3   signature   S45838 ref NP_079359.1  hypothetical protein   FLJ21128 [Homo sapiens]   S45838 ref NP_079359.1  hypothetical protein   FLJ21128 [Homo sapiens]   S5E-49   signature   S458303 pir JC5348 cdd4 protein - Clostridium difficile emb CAA63569.1  (X92982) cdd4 [Clostridium difficile]   0.28   signature   sig	1696	13569915		4.4
Guillardia theta] gb AAF24011.1 AF083031 8     13812078 (AF083031) hypothetical protein [Guillardia theta]   5.2     gi 462193 sp Q06003 GOLI_DROME GOLIATH     PROTEIN (GI PROTEIN) pir  C1495 regulatory     protein G1 - fruit fly (Drosophila melanogaster)     gb AA28582.1  (M97204) goliath protein [Drosophila melanogaster]   9.3     gi 13376638 reflNP_079359.1  hypothetical protein     FIJ21128 [Homo sapiens] db  BAB15001.1      1702   13376638 (AK024781) unnamed protein product [Homo sapiens]   5E-49     gi 7445803 pir  C5348 edd4 protein - Clostridium     difficile   mb CAA63569.1  (X92982) edd4 [Clostridium     difficile   0.28     gi 10175016 db  BAB06115.1  (AP001515)     10175016   BH2396-unknown [Bacillus halodurans]   9     gi 3953531 db  BAA34722.1  (AB015856) ATF6				
gi 462193 sp Q06003 GOLI_DROME GOLIATH   PROTEIN (GI PROTEIN) GI] PROTEIN (GI PROTEIN) priplic 1495 regulatory protein GI - fmit fly (Droscophia melanegaster)   gb AAA28582.1  (M97204) goliath protein [Drosophila melanogaster]   9.3   gi 13376638 ref NP_079359.1  hypothetical protein   FLJ21128 [Homo sapiens] db]BAB15001.1    1702   13376638   (AROZ4781) unamend protein protein (Homo sapiens)   5E-49   gi 7445803 pir  JC5348 cdd4 protein - Clostridium difficile emb CAA63569.1  (X92982) cdd4 [Clostridium difficile]   0.28   gi 10175016 db]BAB06115.1  (AP001515)   0.28   gi 3053531 db]BAA34722.1  (AB015856) ATF6   1706   10175016   BH2396-unknown [Bacillus halodurans]   9   gi 3953531 db]BAA34722.1  (AB015856) ATF6   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   1707   170				
PROTEIN (G1 PROTEIN) pir  IC1495 regulatory   protein G1 - fruit fly (Drosophila melanogaster)   gb AA28582.1  (M97204) goliath protein [Drosophila melanogaster]   9.3	1697	13812078	(AF083031) hypothetical protein [Guillardia theta]	5.2
protein G1 - fruit fly (Drosophila melanogaster)   gjhAAA28582.1](M97204) goliath protein [Drosophila melanogaster]   9,3			gi 462193 sp Q06003 GOLI DROME GOLIATH	
gb AAA28582.1  (M97204) goliath protein [Drosophila melanogaster]   9,3	1			
1699   462193   melanogaster]   9.3				
gij13376638 rd NP_079359.1  hypothetical protein FLJ21128 [Homo sapiens] dhj BAB15001.1  1702 13376638 (AK024781) unnamed protein product [Homo sapiens] 5E-49 gij7445803 pir] JC5348 cdd4 protein - Clostridium difficile emb CAA63569.1  (X92982) cdd4 [Clostridium difficile] 0.28 gij10175016 dbj BAB06115.1  (AP001515) 1706 10175016   BH2396-unknown [Bacillus halodurans] 9 gij3953531 dbj BAA34722.1  (AB015856) ATF6				
FLJ21128 [Homo sapiens] dbj[BAB15001.1]	1699	462193	melanogaster]	9.3
FLJ21128 [Homo sapiens] dbj[BAB15001.1]			L <u></u>	
1702   13376638 (AK024781) unnamed protein product [Homo sapiens]   5E-49				
gi/7445803 pir  JC5348 cdd4 protein - Clostridium difficile emb[CAA65569.1] (X92982) cdd4 [Clostridium difficile mb]   0.28   gi  10175016 dbj  BAB06115.1] (AP001515)   0.28   gi  10175016 dbj  BAB06115.1] (AP001515)   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28   0.28	1700	12276620		5E 40
difficile emb CAA63569.1  (X92982) cdd4 [Clostridium   1705   7445803   difficile]   0.28	1702	13370038		3E-49
1705   7445803   difficile    0.28				
gi 10175016 dbj BAB06115.1  (AP001515)   10175016 BH2396-unknown [Bacillus halodurans] 9   gi 3953531 dbj BAA34722.1  (AB015856) ATF6	1705	7445803		0.28
1706   10175016   BH2396~unknown [Bacillus halodurans]   9   gi 3953531 dbj BAA34722.1  (AB015856) ATF6	1,05	111000		0.20
gi 3953531 dbj BAA34722.1  (AB015856) ATF6	1706	10175016		9
	1709	3953531	[Homo sapiens]	9.1

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 14329676 emb CAC40651.1  (AJ242782) centaurin	
1712	14329676	beta [Homo sapiens]	0.00003
		gi 10432382 emb CAC10340.1  (AL139421) dJ717I23.1	
		(novel protein similar to Xenopus laevis Sojo protein)	
1714	10432382	[Homo sapiens]	0,00006
		gi 8247357 emb CAB92957.1  (AJ401026) hypothetical	
1717	8247357	protein [Thermoanaerobacter thermohydrosulfuricus]	1.4
		gi 7487043 pir  T08926 hypothetical protein T15N24.50	
		- Arabidopsis thaliana emb CAB77061.1  (AL078465)	
		putative protein [Arabidopsis thaliana]	
	i	emb CAB79515.1  (AL161565) putative protein	
1719	7487043	[Arabidopsis thaliana]	0.39
		gi 13507765 ref NP_109714.1  similar to GTPases	
		[Mycoplasma pneumoniae] sp[P75088 Y024_MYCPN	
		PROBABLE GTP-BINDING PROTEIN MG024	
		HOMOLOG (B01_ORF362) pir  S73454 probable GTP	
		binding protein yyaF - Mycoplasma pneumoniae (strain	
	l	ATCC 29342) gb AAB95776.1  (AE000015) similar to	
1721	13507765		6
1505	11051500	gi 11071788 emb CAC14632.1  (AL449144)	
1725	110/1788	hypothetical protein P214.26 [Leishmania major]	9
1.707	14225505	gi 14325595 dbj BAB60498.1  (AP000996) hypothetical	0.5
1727	14325595	protein [Thermoplasma volcanium]	2.5

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
		gi[6498442 dbj[BAA87845.1] (AP000815) ESTs AU081301(E20138),C99280(E10593) correspond to a region of the predicted gene.—Similar to Arabidopsis thaliana chromosome II BAC T26B15 genomic sequence,unknown protein. (Ac004681) [Oryza sativa] dbj[BAB00648.2] (AP002804) ESTs AU081301(E20138),C99280(E10593) correspond to a region of the predicted gene.—Similar to Arabidopsis thaliana chromosome II BAC T26B15 genomic sequence; unknown protein (Ac004681) [Oryza sativa] dbj[BAB17744.1] (AP002862) contains ESTs AU081301(E20138),C99280(E10593)—similar to Arabidopsis thaliana chromosome 2, F12C20.11—unknown protein [Oryza sativa] dbj[BAB44118.1] (AP003103) contains ESTs AU081301(E20138),C99280(E10593)—similar to Arabidopsis thaliana chromosome 2,	TVALUE
1728	6498442		1.1
		gijl11467083 ref[NP_042559.1  NADH dehydrogenase, subunit 3 [Acanthamoeba castellanii] spiQ37382 NU3M_ACACA NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 3 pir  S53860 NADH dehydrogenase (ubiquinone) (EC 1.6.3.3) chain 3 - Acanthamoeba castellanii mitochondrion gb AAD11852.1  (U12386) NADH dehydrogenase,	
1741	11467083	subunit 3 [Acanthamoeba castellanii] gi 12843826 dbi BAB26128.1  (AK009187) putative	5
1742	12843826	[Mus musculus]	0.00000002
1743	4928550	gi 4928550 gb AAD33637.1  (AF133876) DBL alpha protein [Plasmodium falciparum]	6.2
1744	6503033	gi 6503033 gb AAF14557.1 AF176666_1 (AF176666) F-box leucine-rich repeat protein 5 [Xenopus laevis]	3.8
1745	6562750	gi 6562750 cmb CAB62889.1  (AL035475) hypothetical protein, MAL4P2.48 [Plasmodium falciparum]	8.9
1750	7498998	gi 7498998 pir  T16057 hypothetical protein F13D11.2 - Caenorhabditis elegans	5.4
1751	6939792	gi 6939792 dbj BAA90658;1  (AB037920) HA-17 [Clostridium botulinum]	7.3

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 14195008 sp Q9Л55 PLE1_CRIGR PLECTIN 1	
	ļ	(PLTN) (PCN) (300-KDA INTERMEDIATE	
		FILAMENT-ASSOCIATED PROTEIN) (IFAP300)	
		gb AAF70372.1  (AF260753) plectin [Cricetulus	
1754	14195008		1.6
		gi 74519 pir  GNWVR4 structural polyprotein - rubella	
1756	74519	virus (strain Therien)	4.4
		gi 6324076 ref NP_014146.1  Ynl253wp	
		[Saccharomyces cerevisiae] sp P53851 YNZ3_YEAST	
		HYPOTHETICAL 47.2 KD PROTEIN IN SIP3-	
		MRPL30 INTERGENIC REGION pir S63226	
		hypothetical protein YNL253w - yeast (Saccharomyces	
		cerevisiae) emb CAA65491.1  (X96722) ORF N0860	
		[Saccharomyces cerevisiae] emb CAA96160.1	-
1760	6324076	(Z71529) ORF YNL253w [Saccharomyces cerevisiae]	3.6
		gi 1708082 sp P50900 GUX2_CLOSR	
		EXOGLUCANASE II PRECURSOR	
		(EXOCELLOBIOHYDROLASE II) (1,4-BETA-	
	}	CELLOBIOHYDROLASE II) (AVICELASE II)	0
		emb CAA93280.1  (Z69359) avicelase II [Clostridium	
1762	1708082	stercorarium]	1.9
		gi 2224838 emb CAA45388.1  (X63974) putative	
1765	2224838	protein-tyrosine phosphatase [Rhodobacter capsulatus]	5.2
		gi 6513773 gb AAF14748,1 AF197756 1 (AF197756)	
1768	6513773	maturase [Hedvosmum arborescens]	0.63
		gi 14732870 ref XP 029018.1  hypothetical protein	
		FLJ10647 [Homo sapiens] ref[XP 001969.2]	
1775	14732870	hypothetical protein FLJ10647 [Homo sapiens]	1.3
		gi 7521942 pir  T29096 gag polyprotein - murine	
		endogenous retrovirus ERV-L emb CAA73250.1	
1779	7521942	(Y12713) Gag polyprotein [Mus musculus]	0.071
		- 17509 4091-1-17795251 1	
		gi 7508408 pir  T25251 hypothetical protein T24H10.4 - Caenorhabditis elegans emb CAA90944.1  (Z54216)	
1785	7508408	T24H10.4 [Cacnorhabditis elegans]	2.9
1/03	1300408	gil7460247 pir  B71612 hypothetical protein PFB0555c -	2.9
		malaria parasite (Plasmodium falciparum)	
		gb AAC71900.1  (AB001402) hypothetical protein	
1788	7460247	[Plasmodium falciparum]	6.4
1,00	7-100247	It manoman morphum	0.4

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WC0214500 [flig ///E /WO0214500 opc]

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO_	N	DESCRIPTION	P VALUE
		gi 4337102 gb AAD18078.1 AAD18078 (AF129756)	
1789	4337102	G6f [Homo sapiens]	2E-24
		gi 1723494 sp Q10413 YD88_SCHPO VERY	
		HYPOTHETICAL 13.3 KD PROTEIN C1F3.08C IN	
		CHROMOSOME I pir T38079 very hypothetical	
		protein SPAC1F3.08c - fission yeast	
		(Schizosaccharomyces pombe) emb CAA94626.1	
		(Z70690) very hypothetical protein	
1792	1723494	[Schizosaccharomyces pombe]	7.2
		gi 4589921 dbj BAA76927.1  (AB017192)	
		molybdopterin biosynthesis protein [Clostridium	
1793	4589921	perfringens]	4.2
		gi 7494302 pir  E71611 hypothetical protein PFB0580w -	
		malaria parasite (Plasmodium falciparum)	
		gb AAC71905.1  (AE001404) hypothetical protein	
1799	7494302	[Plasmodium falciparum]	3.3
		gi 14318508 ref NP 116641.1  Ies1p [Saccharomyces	
		cerevisiael splP43579 YFB3 YEAST	
		HYPOTHETICAL 78.8 KD PROTEIN IN HSP12-	
		HXT10 INTERGENIC REGION pir S48316 probable	
		membrane protein YFL013c - yeast (Saccharomyces	
		cerevisiae) emb CAA86347.1  (Z46255) orf, len: 692,	
		CAI: 0.14 [Saccharomyces cerevisiae] dbj BAA09225.1	
1800	14318508	(D50617) YFL013C [Saccharomyces cerevisiae]	6.6
		gi 14325595 dbj BAB60498.1  (AP000996) hypothetical	
1802	14325595	protein [Thermoplasma volcanium]	2
		gi 7520399 pir  T11689 NADH dehydrogenase	· ·
		(ubiquinone) (EC 1.6.5.3) chain 5 - Graphium sarpedon	
		mitochondrion (fragment) dbj BAA28187.1	
		(AB013147) NADH dehydrogenase subunit 5	
1809	7520399	[Graphium sarpedon]	4.9
		gi 423981 pir  A46193 88K E-26-specific domain	
	ļ	protein Pok - fruit fly (Drosophila melanogaster)	
		dbj BAA01080.1  (D10228) Ets domain protein	
1816	423981	[Drosophila melanogaster]	0.23
		gi 7463036 pir  C70177 beta-glucosidase homolog -	
		Lyme disease spirochete gb AAC66976.1  (AE001163)	
1818	7463036		8.4
		gi 3851471 gb AAC72292.1  (AF037295) sulfonylurea	
1819	3851471	receptor-1 [Mus musculus]	3.4

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 9845030 dbj BAB11914.1  (AB034726) 5-	
1820	9845030	oxoprolinase precursor [Alcaligenes faecalis]	5.8
		gi 6599044 emb CAB63584.1  (AJ251826) BMP2/4	
1821	6599044	protein [Asterias rubens]	6.7
		gi 7715984 gb AAF68235,1 AF206244 1 (AF206244)	
1822	7715984	seroreactive antigen BMN1-2 [Babesia microti]	0,067
		gi 7293329 gb AAF48709.1  (AE003504) CG5162 gene	
1823	7293329	product [Drosophila melanogaster]	2.8
		gi 2136095 pir  I58381 receptor tyrosine kinase isoform	
		FLT4 long - human (fragment) gblAAB28539.1	
		(S66407) receptor tyrosine kinase isoform FLT4 long.	
		FLT41 {C-terminal} [human, Peptide Partial, 71 aa]	
1824	2136095		0.0007
		gi 7522093 pir  T30809 plasminogen related growth	
		factor receptor 3 - Fugu rubripes emb CAA09086.1	
		(AJ010317) plasminogen related growth factor receptor	
1826	7522093	3 [Takifugu rubripes]	4.7
		gi 13622999 gb AAK34670.1  (AE006621) hypothetical	
1832	13622999	protein [Streptococcus pyogenes M1 GAS]	3.7
		gi 6319504 ref NP_009586.1  Ybr030wp	
		[Saccharomyces cerevisiae] sp P38222 YBO0_YEAST	
		HYPOTHETICAL 62.6 KD PROTEIN IN CDS1-	
		RPL4A INTERGENIC REGION pir  S45886	
		hypothetical protein YBR030w - yeast (Saccharomyces	
		cerevisiae) emb CAA53686.1  (X76078) YBR0314	
		[Saccharomyces cerevisiae] emb CAA84972.1	
		(Z35899) ORF YBR030w [Saccharomyces cerevisiae]	
		prf  2206497K ORF YBR0314 [Saccharomyces	
1834	6319504		0.65
		gi 7290986 gb AAF46425.1  (AE003445) CG3898 gene	
1835	7290986	product [Drosophila melanogaster]	3.1
		gi 14725330 ref XP_002254.2  mitochondrial	
	l	translational initiation factor 2 precursor [Homo	
1840	14725330	sapiens]	4.2

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	_ N	DESCRIPTION	P VALUE
		gi 7509362 pir  T31492 hypothetical protein	
		Y116A8C.20 - Caenorhabditis elegans	
		emb CAB55126.1  (AL117204) predicted using	
		Genefinder-contains similarity to Pfam domain:	
		PF00642 (Zinc finger C-x8-C-x5-C-x3-H type (and	
		similar).), Score=71.0, E-value=3.1e-19, N=2~cDNA	
		EST yk247a8.5 comes from this gene [Caenorhabditis	
1842	7509362	elegans]	8.4
		gi 6093793 sp Q64181 PROP_CAVPO PROPERDIN	
		PRECURSOR gb AAB35918.1  (S81116) properdin	
1846	6093793	[guinea pigs, spleen, Peptide, 470 aa] [Cavia]	4.1
		gi 7300538 gb AAF55691.1  (AE003727) CG7411 gene	
1848	7300538	product [Drosophila melanogaster]	2.7
		gi 1730077 sp P18160 KYK1 DICDI NON-	
		RECEPTOR TYROSINE KINASE SPORE LYSIS A	
		(TYROSINE-PROTEIN KINASE 1) pir  T18276 non-	
		receptor tyrosine kinase - slime mold (Dictyostelium	
		discoideum) gb[AAB41125.1] (U32174) non-receptor	
1849	1730077	tyrosine kinase [Dictyostelium discoideum]	8.4
1015	1750077	gi 5326919 emb CAB46239.1  (AJ133488) SCO-	
1851	5326919	spondin [Bos taurus]	2.7
		gi 13811938 emb CAC03433.2  (AL118505)	
		dJ1056H1.2.1 (novel protein similar to mitogen	
1853	13811938	inducible protein MIG-2 (isoform 1)) [Homo sapiens]	2E-21
		1005 to 771 (DVD 0502 (5 1))	
		gi 8954377 ref[NP_059365.1  haem lyase  Cyanidioschyzon merolae  pir A58932 cytochrome C-	
		type biogenesis protein CCMF - Cyanidioschyzon	
		merolae mitochondrion dbj BAA36527.1  (D89861)	
		cytochrome C-type biogenesis protein CCMF	
1859	8954377	[Cyanidioschyzon merolae]	2.3
	12.277	L-J-	
		gi 13376638 ref[NP_079359.1  hypothetical protein	
		FLJ21128 [Homo sapiens] dbj BAB15001.1	
1860	13376638	(AK024781) unnamed protein product [Homo sapiens]	2E-47
		gi 14602664 gb AAH09855.1 AAH09855 (BC009855)	
1862	14602664	Similar to nucleolin [Homo sapiens]	4E-45

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 7512078 pir  T30879 dynein heavy chain isotype 5A -	
		sea urchin (Tripneustes gratilla) (fragment)	
i		gb AAA63591.1  (U03977) dynein heavy chain isotype	
1864	7512078	5A [Tripneustes gratilla]	1.7
		gi 7688347 emb CAB89836.1  (AJ242516) AmpE	
1865	7688347	protein [Salmonella typhimurium]	2.8
		gi 4758568 ref NP_004497.1  heat shock transcription	
		factor 2 [Homo sapiens] ref[XP_004466.2] heat shock	
		transcription factor 2 [Homo sapiens] ref[XP_051573.1]	
		heat shock transcription factor 2 [Homo sapiens]	
		sp Q03933 HSF2_HUMAN HEAT SHOCK FACTOR	
		PROTEIN 2 (HSF 2) (HEAT SHOCK	
		TRANSCRIPTION FACTOR 2) (HSTF 2) pir  A41138	
		heat shock transcription factor HSF2 - human	
1867	4758568	gb AAA36017.1  (M65217) HSF2 [Homo sapiens]	0.33
1869	13358509	gi 13358509 ref NP_078688.1  orf107 [lymphocystis disease virus 1]	8.1
1869	13338309		0.1
		gi 6580323 emb CAB63392.1  (AL132864) cDNA EST	
		EMBL:T00048 comes from this gene~cDNA EST EMBL:T00047 comes from this gene~cDNA EST	
ļ		yk390e6.3 comes from this gene~cDNA EST yk512a3.3	
		comes from this gene~cDNA EST yk512a3.5 comes	
	Į	from this gene~cDNA EST yk532a2.3 comes from this	
1870	6580323	ge>	0.012
1070	0000022	gi 11359776 pir T45059 hypothetical protein	
		Y39B6B.gg [imported] - Caenorhabditis elegans	
		emb CAB60938.1  (AL132896) predicted using	
		Genefinder; preliminary prediction [Caenorhabditis	
1871	11359776		3.6
		gi 5453171 gb AAD43464.1  (AF113915) pre-mRNA	
1876	5453171	splicing factor [Heterodera glycines]	2.1
		gi 5453171 gb AAD43464.1  (AF113915) pre-mRNA	
1877	5453171	splicing factor [Heterodera glycines]	2.1
		gi 14752353 ref XP_050519.1  annexin A2 [Homo	
1880	14752353	sapiens]	0.27
		gi 7494821 pir  T31996 hypothetical protein B0281.5 -	
		Caenorhabditis elegans gb AAB66084.1  (AF016666)	
		similar to human tumor necrosis factor-alpha-induced	
1881	7494821	protein B12 (NID:g179304) [Caenorhabditis elegans]	3.8

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
_NO	N	DESCRIPTION	P VALUE
		- 14904931 : HG2C052 - 4 1 : 14 - GEG	
		gi 480482 pir  S36953 cytochrome-c oxidase (EC	
		1.9.3.1) chain III - Herpetomonas samuelpessoai	
		mitochondrion gb AAD09166.1  (L10852) cytochrome	
1883	480482	oxidase subunit III [Herpetomonas pessoai]	0.78
		gi 5174493 ref NP_006050.1  laminin, gamma 3	
		precursor [Homo sapiens]	
		gb AAD36991.1 AF041835_1 (AF041835) laminin	
1885	5174493	gamma 3 chain precursor [Homo sapiens]	3E-12
	1	gi 8922792 ref NP_060753.1  hypothetical protein	
		FLJ10956 [Homo sapiens] ref[XP_007214.2]	
		hypothetical protein FLJ10956 [Homo sapiens]	
		dbj BAA91925.1  (AK001818) unnamed protein product	
1886	8922792	[Homo sapiens]	5E-39
	ŀ	gi 7479561 pir  T35135 hypothetical protein SC4H8.04c	
		SC4H8.04c - Streptomyces coelicolor	
		emb CAA15871.1  (AL020958) hypothetical protein	
1889	7479561	SC4H8.04c [Streptomyces coelicolor A3(2)]	0.5
		gi 7461128 pir  T03057 hypothetical protein 032R -	
		Chilo iridescent virus gb AAB94431.1  (AF003534)	
1890	7461128	hypothetical protein 032R [Chilo iridescent virus]	0.33
		gi 8778367 gb AAF79375.1 AC007887_34 (AC007887)	
1891	8778367	F15O4.28 [Arabidopsis thaliana]	6.6
		gi 13475247 ref NP_106811.1  unknown protein	
		[Mesorhizobium loti] dbj BAB52597.1  (AP003008)	
1894	13475247	unknown protein [Mesorhizobium loti]	2.3
		gi 2285958 emb CAA70903.1  (Y09763) GABRE	
1895	2285958	[Homo sapiens]	2.7
		gi 14729939 ref XP_038475.1  DKFZP564J102 protein	
1896	14729939	[Homo sapiens]	0.007
	1		
		gi 7504499 pir  T32750 hypothetical protein F57B10.1 -	
		Caenorhabditis elegans gb AAB96719.1  (AF039713)	
1898	7504499	Hypothetical protein F57B10.1 [Caenorhabditis elegans]	4.2
		gi 10581310 gb AAG20067.1  (AE005086) methionyl	
1900	10581310	aminopeptidase; Map [Halobacterium sp. NRC-1]	8.9
		gi 111814 pir  S21347 hypothetical protein 3 - rat	
1902	111814	emb CAA37646.1  (X53581) ORF3 [Rattus norvegicus]	0,3
		gi 7290766 gb AAF46211.1  (AE003439) CG4557 gene	
1903	7290766	product [Drosophila melanogaster]	0.38

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS N	DVGCD-DWC011	
NO	N	DESCRIPTION	P VALUE
		"IS 1205211 LIGADASS CO. 11 (A YOURGON COMP.)	
		gi 5139521 cmb CAB45562.1  (AJ238798) CTRP protein [Plasmodium berghei] dbi BAA82322.1	
		(AB027129) adhesive protein-like molecule	
		[Plasmodium berghei] gb AAF73158.1 AF149771 1	
1904	5120521	(AF149771) ookinete protein [Plasmodium berghei]	8
1904	3139321	gi[7489900]pir[T18287 protein-tyrosine kinase (EC	
		2.7.1.112) - slime mold (Dictyostelium discoideum)	
		gb[AAB04999.1] (U64830) protein tyrosine kinase	
1905	7489900	[Dictyostelium discoideum]	5.3
1903	7489900	gi 6002776 gb AAF00134.1 AF149806 1 (AF149806)	3.3
1908	6002776	hypothetical protein [Oryza sativa]	0.15
1900	0002770	nypomonoai protem [Oryza sauva]	0.13
1913	14773348	gi 14773348 ref XP 038450.1  20849 [Homo sapiens]	2E-50
1715	11770010	gi 7301187 gb AAF56319.1  (AE003748) CG5794 gene	
1916	7301187	product [Drosophila melanogaster]	8
1710	7501107	gi 3378685 emb CAA76071.1  (Y16104) replicase	
1918	3378685	protein [Physalis mottle tymovirus]	0.13
		gi 4501915 ref NP 003807.1  a disintegrin and	
		metalloproteinase domain 9 preproprotein; meltrin	
		gamma [Homo sapiens] gb[AAC50403.1] (U41766)	
		metalloprotease/disintegrin/cysteine-rich protein	
1919	4501915		0.002
		gi 14587070 gb AAK70463.1 AF387344_4 (AF387344)	
1922	14587070	spore germination protein GerLC [Bacillus cereus]	8.4
		gi 7291161 gb AAF46595.1  (AE003450) CG2892 gene	
1924	7291161	product [Drosophila melanogaster]	6
		gi 7446016 pir  E70895 hypothetical glycine-rich protein	
		Rv1087 - Mycobacterium tuberculosis (strain H37RV)	
		emb CAA17203.1  (AL021897) PE_PGRS	
1925	7446016	[Mycobacterium tuberculosis]	3.6
		gi 11595522 cmb CAC18316.1  (AL451022)	
1926	11595522	hypothetical protein [Neurospora crassa]	5.9
		gi 14043326 gb AAH07658.1 AAH07658 (BC007658)	
1930	14043326	Unknown (protein for MGC:747) [Homo sapiens]	8E-76
		gi 13810543 dbj BAB43950.1  (AB051633) ookinete	۱
1931	13810543	surface protein Pos28-2 [Plasmodium ovale]	3.1
		gi 7206826 gb AAF39985.1  (AC006696) contains	-
	=======================================	similarity to other proline-rich proteins [Caenorhabditis	
1933	7206826	[elegans]	6.6

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS	PHACHAMAN		
NO	N	DESCRIPTION	P VALUE	
		gi 14749721 ref XP_027893.1  similar to ALU		
1004	14540501	SUBFAMILY SBI SEQUENCE CONTAMINATION		
1934	14749721	WARNING ENTRY (H. sapiens) [Homo sapiens]	2.1	
		gi 12324211 gb AAG52077.1 AC012679_15		
		(AC012679) putative proline-rich protein precursor;		
1935	12324211	93710-91881 [Arabidopsis thaliana]	2.1	
		gi 4204305 gb AAD10686.1  (AC003027) Hypothetical		
1940	4204305	protein [Arabidopsis thaliana]	5.4	
		gi 14783118 ref XP_043478.1  hypothetical protein		
1955	14783118	XP_043478 [Homo sapiens]	4.7	
l		gi 13375860 ref NP_078907.1  hypothetical protein		
l		FLJ23342 [Homo sapiens] dbj BAB15618.1		
1959	13375860	(AK026995) unnamed protein product [Homo sapiens]	0.2	
	,	gi 7304316 gb AAF59348.1  (AE003844) CG2052 gene		
1962	7304316	product [Drosophila melanogaster]	8.2	
		gi 7513579 pir  T09064 1-acylglycerol-3-phosphate O-		
1		acyltransferase (EC 2.3.1.51) - mouse gb AAB82009.1		
		(AF030001) lysophatidic acid acyl transferase-alpha		
1963	7513579	[Mus musculus]	1.1	
		gi 7512671 pir  T12545 hypothetical protein		
ĺ		DKFZp434N074.1 - human (fragments)		
		emb CAB46377.1  (AL096732) hypothetical protein		
1964	7512671	[Homo sapiens]	3.1	
		gi 13249541 gb AAK15414.1  (AY015597)		
		dissimilatory sulfite reductase subunit B [uncultured		
1972	13249541	sulfate-reducing bacterium]	2	
		gi 8134766 sp Q9ZES2 TRPE_BUCTC		
		ANTHRANILATE SYNTHASE COMPONENT I		
		emb CAA09993.1  (AJ012333) anthranilate synthase		
1976	8134766	large subunit [Buchnera aphidicola]	5.6	
		gi 7518768 pir  A71111 hypothetical protein PH0656 -		
		Pyrococcus horikoshii dbj BAA29747.1  (AP000003)		
1977	7518768	107aa long hypothetical protein [Pyrococcus horikoshii]	9.3	
1		n		
		gi 14749721 ref XP_027893.1  similar to ALU		
1,000	1474070	SUBFAMILY SB1 SEQUENCE CONTAMINATION	1.0	
1982	14/49/21	WARNING ENTRY (H. sapiens) [Homo sapiens]	1.9	

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	PHICHMAN	
NO_	N	DESCRIPTION	P VALUE
		gi 2498123 sp O02833 ALS_PAPHA INSULIN-LIKE	
		GROWTH FACTOR BINDING PROTEIN	
		COMPLEX ACID LABILE CHAIN PRECURSOR	
		(ALS) pir  JC5239 insulin-like growth factor acid-labile	W.
1983	2498123		4.4
		gi 2911545 emb CAA75449,1  (Y15173) E6 protein	
1990	2911545	[Human papillomavirus type 75]	4.2
		gi 1090764 prf  2019432A cyclin-dependent kinase 5	
1992	1090764	activator [Bos taurus]	2.1
		gi 7446170 pir  T02529 myb-related protein F13M22.13	
		- Arabidopsis thaliana gb AAC23633.1  (AC004684)	
		putative MYB family transcription factor [Arabidopsis	
		thaliana] gb AAD53101.1 AF175996_1 (AF175996)	
1993	7446170	putative transcription factor [Arabidopsis thaliana]	5.9
		gi 4501915 ref NP_003807.1  a disintegrin and	
		metalloproteinase domain 9 preproprotein; meltrin	
	i	gamma [Homo sapiens] gb AAC50403.1  (U41766)	
		metalloprotease/disintegrin/cysteine-rich protein	
1994	4501915	precursor [Homo sapiens]	0.002
		gi 12853260 dbj BAB29697.1  (AK015063) putative	
1997	12853260	[Mus musculus]	6.1
		gi 6492289 gb AAF14258.1 AF137068_1 (AF137068)	
2000	6492289		4.6
		gi 2828501 sp P40899 ISP3_SCHPO SEXUAL	
		DIFFERENTIATION PROCESS PROTEIN ISP3	
		pir  T38112 sexual differentiation process protein isp3 -	
		fission yeast (Schizosaccharomyces pombe)	
		emb CAB03599.1  (Z81312) sexual differentiation	
		process protein isp3; meiotic expression upregulated	
2001	2828501	[Schizosaccharomyces pombe]	7.9
		gi 4206157 gb AAD11433.1  (AF109404) transposase	
2022	4206157	[Streptomyces scabiei]	3.4
		-1114CC224 FNID 0C2847 1  ODE1 ' 4 :	
	Ì	gi]11466224 ref[NP_062847.1  ORF1, contains 4 trans	
		membrane regions, putative [Physarum polycephalum]	
2026	11466334	dbj BAB08081.1  (AB027295) ORF1, contains 4 trans	1.2
2026	11406224	membrane regions, putative [Physarum polycephalum]	1.2

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	t Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	DIVITION
NO	IN	DESCRIPTION	P VALUE
		-17507526-1-177247201	
ł	ł	gi 7507536 pir  T24739 hypothetical protein T09E11.4 - Caenorhabditis elegans emb CAB03533.1  (Z81147)	ł
		contains similarity to Pfam domain: PF01757 (Domain	
		of unknown function), Score=594.0, E-value=3e-175,	
2027	7507536		4.3
2021	7307330	gi 6102749 emb CAB59307,1  (AJ236287) NADH	4.3
2028	6102749	dehydrogenase subunit F [Carphalea glaucescens]	1.9
2020	0102743	gi 7486224 pir  T08553 hypothetical protein	1.9
		F27B13.160 - Arabidopsis thaliana emb CAB43667.1	
		(AL050352) putative protein [Arabidopsis thaliana]	
		emb CAB79750.1  (AL161575) putative protein	
2031	7486224	[Arabidopsis thaliana]	2.7
2031	7400224	gi 7462619 pir F72210 hypothetical protein TM1801 -	2.1
		Thermotoga maritima (strain MSB8)	
		gb AAD36864.1 AE001817 11 (AE001817)	
2032	7462619	hypothetical protein [Thermotoga maritima]	6.2
2032	7402015	gi 7296128 gb AAF51422.1  (AE003587) CG4629 gene	0.2
2035	7296128	product [Drosophila melanogaster]	3.4
2000	7250120	product [D1000pmin inchanogustor]	3.4
		gi 14760974 ref XP 034809.1  similar to putative gag-	
2038	14760974	pro-pol polyprotein (H. sapiens) [Homo sapiens]	1E-13
		gi 14601134 ref NP 147662.1  hypothetical protein	12.10
		[Aeropyrum pernix] pir  H72698 hypothetical protein	
		APE1008 - Aeropyrum pernix (strain K1)	
		dbj BAA79992.1  (AP000060) 123aa long hypothetical	
2041	14601134	protein [Aeropyrum pernix]	2.6
		-14507527  (NID 002000 II - 1 - 1 - 1 - 1 - 1	
		gi 4507537 ref NP_003260.1  nuclear receptor subfamily	
		2, group E, member 1; tailless (Drosophila) homolog; tailless homolog (Drosophila) [Homo sapiens]	
		ref[XP 004530.1] 54551 [Homo sapiens]	
		ref[XP 038737.1] nuclear receptor subfamily 2, group	
		E, member 1 [Homo sapiens]	
		spiO9Y466 NR21 HUMAN ORPHAN NUCLEAR	
		RECEPTOR NR2E1 (NUCLEAR RECEPTOR TLX)	
		(TAILLESS HOMOLOG) (TLL) (HTLL)	
		emb CAA73725.1  (Y13276) Tailless protein [Homo	
		sapiens] emb CAB75626,1  (AL078596) dJ429G5,1	
		(nuclear receptor subfamily 2, group E, member 1)	
		[Homo sapiens] gb AAG31945.1 AF220532 1	
2043	4507537	(AF220532) orphan nuclear receptor [Homo sapiens]	9E-10

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 7300319 gb AAF55480.1  (AE003719) CG7305 gene	
2044	7300319	product [Drosophila melanogaster]	3.3
		gi 7498832 pir [T34212 hypothetical protein F10E7.4 - Caenorhabditis elegans gb AA.82427.1  (U41264) coded for by C. elegans cDNA yk99a6.5; coded for by	
		C. elegans cDNA yk72g6.5; coded for by C. elegans	100
		cDNA yk99a6.3; coded for by C. elegans cDNA	
2046	# 400000	yk72g6.3; coded for by C. elegans cDNA yk127a2.5;	
2046	7498832	coded for by C. elegans cDNA yk127a2.3; Simila>	9.7
		gil 13643847heflXP_011044.2l 8-oxoguanine DNA glycosylase [Homo sapiens] reflXP_016414.1l 8- oxoguanine DNA glycosylase [Homo sapiens] reflXP_016415.1l 8-oxoguanine DNA glycosylase [Homo sapiens] reflXP_031967.1l 8-oxoguanine DNA glycosylase [Homo sapiens] reflXP_031959.1l 8- oxoguanine DNA glycosylase, isoform 2d [Homo	
		sapiens] ref[XP_031961.1  8-oxoguanine DNA	
		glycosylase, isoform 1a [Homo sapiens] ref[XP 052277.1  similar to 8-oxoguanine DNA	
2050	12642947	glycosylase (H. sapiens) [Homo sapiens]	1.2
2030	13043647	grycosyrase (rr. sapiens) [rionio sapiens]	1.2
2051	7474551	gi 7474551 pir  E69792 conserved hypothetical protein yeeA - Bacillus subtilis emb CAB12496.1  (Z99107) similar to hypothetical proteins [Bacillus subtilis]	7
2031	1414331	similar to hypotheticar proteins [Bacillus suotins]	
		gij3913201 sp Q58511 CCA_METJA TRNA NUCLEOTIDYLTRANSFERASE (TRNA ADENYLYLTRANSFERASE) (TRNA CCA- PYROPHOSPHORYLASE) (CCA-ADDING ENZYME) gb AAB99114.1  (U67554) URNA	
2060	3913201	nucleotidyltransferase (cca) [Methanococcus jannaschii]	5.9
2072	13186342	gi 13186342 gb AAK15384.1  (AF211134) valyl-tRNA synthetase [Carsonella ruddii]	2.8

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WC0214560 [fle://E/WO0214500 opc]__

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION .	P VALUE
110		DESCRIPTION .	1 VALUE
	ľ	gi 6225820 sp O04226 P5CS ORYSA DELTA 1-	
	l	PYRROLINE-5-CARBOXYLATE SYNTHETASE	
		(P5CS) [INCLUDES: GLUTAMATE 5-KINASE	
		(GAMMA-GLUTAMYL KINASE) (GK); GAMMA-	
1	ŀ	GLUTAMYL PHOSPHATE REDUCTASE (GPR)	
		(GLUTAMATE-5-SEMIALDEHYDE	
1		DEHYDROGENASE) (GLUTAMYL-GAMMA-	
		SEMIALDEHYDE DEHYDROGENASE)] pir  T03695	
		delta 1 pyrroline-5-carboxylate synthetase - rice	
1		dbj BAA19916.1  (D49714) deltal-pyrroline-5-	
2075	6225820	carboxylate synthetase [Oryza sativa]	9.4
		gi 4028153 gb AAC96117.1  (AF083221) putative	
2077	4028153	neurotransmitter receptor [Takifugu rubripes]	2
i		gi 4902680 emb CAB43550.1  (AL031673) dJ694B14.3	
		(novel haloacid dehalogenase-like hydrolase family	
		protein similar to (archaea) bacterial proteins) [Homo	
2080	4902680	sapiens]	2.9
]		gi 6322760 ref NP_012833.1  Ykl090wp	
ľ		[Saccharomyces cerevisiae] sp P36075 YKJ0_YEAST	
		HYPOTHETICAL 50.9 KD PROTEIN IN BUD2-	
1		MIF2 INTERGENIC REGION pir \$\frac{1}{37915}\$ hypothetical protein YKL090w - yeast (Saccharomyces)	
J		cerevisiae) emb CAA81928.1  (Z28090) ORF	
2084	6322760	YKL090w [Saccharomyces cerevisiae]	2.8
2004	0322100	gi 6635084 emb CAB64573.1  (AL135930) hypothetical	2.0
2085	6635084	protein L4738.02 [Leishmania major]	3.6
2005	0055001	protein 27730.02 [Dominante major]	
		gi 7487726 pir  T05814 hypothetical protein T5K18.90 -	
1		Arabidopsis thaliana emblCAA18618,1 (AL022580)	
		hypothetical protein [Arabidopsis thaliana]	
1		emb CAB78933.1  (AL161550) hypothetical protein	
2086	7487726	[Arabidopsis thaliana]	5.9
		gi 13385468 ref[NP_080247.1  RIKEN cDNA	
		2900001A12 gene [Mus musculus] dbj BAB28377.1	
1		(AK012645) putative [Mus musculus] dbj BAB28865.1	
2089	13385468	(AK013457) putative [Mus musculus]	4E-11
		gi 14742770 ref XP_039393.1  KIAA1550 protein	
2090	14742770	[Homo sapiens]	3.2
		gi 2982251 gb AAC32113.1  (AF051208) putative RNA-	
2092	2982251	binding protein [Picea mariana]	7.6

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
l !		gi 12718478 emb CAC28807.1  (AL513466)	
2095	12718478	hypothetical protein [Neurospora crassa]	3.1
		gi 5532964 gb AAD44957.1  (AF156655) MHC class I	
2096	5532964	heavy chain [Ambystoma mexicanum]	9.3
1		gi 1711658 sp P54797 T10_MOUSE SER/THR-RICH	
		PROTEIN T10 IN DGCR REGION pir   S37488 gene	
2097	1711650	T10 protein - mouse emb CAA52612.1  (X74504) T10 [Mus musculus]	4.0
2097	1711658	gi 14193393 gb AAK55953.1 AF268062 2 (AF268062)	4.3
Į.		RNA polymerase beta-prime subunit [Candidatus	
2098	14102202	Carsonella ruddii]	2.1
2098	14193393	gi 7243081 dbi BAA92588.1  (AB037771) KIAA1350	2.1
2099	7243081	protein [Homo sapiens]	7E-97
2099	7243001	· · · · · · · · · · · · · · · · · · ·	/E-9/
		gi 11357181 pir  T49996 AtAGP4 - Arabidopsis	
		thaliana gb AAC77826.1  (AF082301) arabinogalactan- protein [Arabidopsis thaliana]	
		gb AAD38870.1 AF060874 1 (AF060874) AtAGP4	
		[Arabidopsis thaliana] emb[CAB89400.1] (AL353995)	
		AtAGP4 [Arabidopsis thaliana]	
1		gb AAK49601.1 AF372885 1 (AF372885)	
		AT5g10430/F12B17 220 [Arabidopsis thaliana]	
1		gb AAK68734.1  (AY042794) AtAGP4 [Arabidopsis	
2100	11357181		1.2
		gi 7497369 pir  T32512 hypothetical protein C44B12.4 -	
		Caenorhabditis elegans gb AAB88327.1  (AF036692)	
1		Hypothetical protein C44B12.4 [Caenorhabditis	
2103	7497369		6.7
		gi 14485227 gb AAK62977.1 AF384372_3 (AF384372)	
2109	14485227	surface antigen [Hepatitis B virus]	5.7
		gi 9711862 dbj BAB07956.1  (AP002524) putative	
		extensin-like protein [Oryza sativa] dbj BAB33013.1	
		(AP003118) putative extensin-like protein [Oryza	
2111	9711862	sativa]	10
		gi 9759203 dbj BAB09740.1  (AB015476) heat shock	
		transcription factor HSF30-like protein [Arabidopsis	
2114	9759203	thaliana]	8.4
		100 40 TOL 114 4 4 4 4 4 0 0 1 0 Theorem 1	
2115	204070	gi 204070 gb AAA41130.1  (M22030) electron transfer	0.75
2115	204070	flavoprotein alpha-subunit [Rattus norvegicus]	0.75
		gi 3638957 gb AAC36301.1  (AC004877) sco-spondin-	
		mucin-like; similar to P98167 (PID:g1711548); details	
2117		of intron/exon structure uncertain [Homo sapiens]	8.1
2111	7020237	or meron exon structure uncertain (110mo sapiens)	0.1

	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE	
		Description 1101	1 VALUE	
		gi 14250436 gb AAH08653.1 AAH08653 (BC008653)		
2121	14250436	Similar to LRP16 protein [Mus musculus]	9.3	
		- 2147C05201 GVP 020700 11 7071 FV		
2126	14760522	gi 14760522 ref XP_038798.1  7371 [Homo sapiens] gb AAK01445.1  (AF334585) NIR3 [Homo sapiens]	5E-39	
	11700000	gi 3024944 sp Q58366 Y956 METJA	315-39	
ļ	l	HYPOTHETICAL PROTEIN MJ0956 pir D64419		
		hypothetical protein MJ0956 - Methanococcus		
ļ		jannaschii gb AAB98969.1  (U67539) M. jannaschii		
2134	2024044	predicted coding region MJ0956 [Methanococcus jannaschii]		
2134	3024944	gi 6606266 gb AAF19148.1 AF158634 1 (AF158634)	8.4	
2135	6606266	Vrgal [Aegilops ventricosa]	7.1	
		gi 1729878 sp P54410 TCPH TETTH T-COMPLEX		
[		PROTEIN 1, ETA SUBUNIT (TCP-1-ETA) (CCT-		
		ETA) pir  S71338 t-complex protein 1 theta chain -		
		Tetrahymena thermophila (fragment) gb AAC47007.1		
		(U46028) CCTeta [Tetrahymena thermophila] prf  2209286B chaperonin CCT-eta [Tetrahymena		
2137	1729878	thermophilal	4.4	
		gi 3688193 emb CAA08995.1  (AJ010091) MAP3K		
2141	3688193	alpha 1 protein kinase [Brassica napus]	6.9	
		<u></u>		
		gi 7491910 pir  T41367 hypothetical protein		
		SPCC4G3.09c - fission yeast (Schizosaccharomyces pombe) emb CAB09776.1  (Z97052) hypothetical		
2142	7491910	protein [Schizosaccharomyces pombe]	3.3	
		gi 6513832 gb AAF14807.1 AF197815 1 (AF197815)		
2144	6513832	maturase [Alisma plantago-aquatica]	1.5	
	ĺ			
		gi 6323084 ref NP_013156.1  transcription factor,		
		probable member of histone acetyltransferase SAGA complex; Spt8p [Saccharomyces cerevisiae]		
		sp P38915 SPT8 YEAST TRANSCRIPTION		
		FACTOR SPT8 pir  S47898 transcription factor SPT8 -		
		yeast (Saccharomyces cerevisiae) gb AAA53585.1		
		(M94955) transcription factor [Saccharomyces		
		cerevisiae] cmb CAA64302.1  (X94607) transcription factor [Saccharomyces cerevisiae] emb CAA97585.1		
2148	6323084	(Z73227) ORF YLR055c [Saccharomyces cerevisiae]	5.8	
		i		

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		Proteins)	
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 1074219 pir  S49239 hypothetical protein 2	
		(capsulation locus) - Hacmophilus influenzae (strain	
2151	1074219	RM107)	4
		gi 10946710 ref NP_067350.1  Rhesus blood group-	
		associated B glycoprotein; Rh type B glycoprotein [Mus	
		musculus] gb AAF19371.1  (AF193808) Rh type B	
2153	10946710	glycoprotein [Mus musculus]	3,5
		gi 9625644 ref NP_039895.1  BDLF2 late reading frame	
		[Human herpesvirus 4] sp P03225 BDL2_EBV	•
		PROTEIN BDLF2 pir QQBE44 BDLF2 protein -	
		human herpesvirus 4 (strain B95-8) emb CAA24836.1	
		(V01555) BDLF2 late reading frame [Human	
2159	9625644	herpesvirus 4]	3.7
		gi 220578 dbj BAA00447.1  (D00570) open reading	
2160	220578	frame (251 AA) [Mus musculus]	4.7
		gi 9633076 ref NP_050182.1  B4 [Human herpesvirus	
		6B] pir  T44148 hypothetical protein B4 [imported] -	
		human herpesvirus 6 (strain Z29)	
		gb AAD49620.1 AF157706_7 (AF157706) B4 [Human	
2165	9633076	herpesvirus 6B]	0.057
		gi 135838 sp P01267 THYG_BOVIN	
		THYROGLOBULIN PRECURSOR pir  UIBO	
		thyroglobulin precursor - bovine emb CAA26584.1	
		(X02815) thyroglobulin precursor [Bos taurus]	
2168	135838	prf  1109240A thyroglobulin [Bos taurus]	2
		gi 11360154 pir  T46337 hypothetical protein	
		DKFZp434O2413.1 - human (fragment)	
		emb CAB70664.1  (AL137265) hypothetical protein	
2173	11360154	[Homo sapiens]	3.7
		gi 181400 gb AAA35748.1  (M34225) cytokeratin 8	
2177	181400	[Homo sapiens]	7E-53
		gi 11347010 pir  B81303 probable membrane protein	
		Cj1013c [imported] - Campylobacter jejuni (strain	
		NCTC 11168) emb CAB73269.1  (AL139077) putative	
2181	11347010	membrane protein [Campylobacter jejuni]	0.1
		gi 8745261 gb AAF78857.1 AF134516_1 (AF134516)	
2182	8745261	VP4 [Banna virus]	6

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# WO 02/14500

WC0214500 [fla //E /WO0214500 opc]

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS	DESCRIPTION	P VALUE
		Discussion Tion	1 TIBOL
		gi 285275 pir  A43963 envelope glycoprotein G(envelope	
		glycoprotein G1, envelope glycoprotein G2) -	
		Hantavirus sp.=Puumala virus gb AAB22506.1	
		envelope glycoprotein G(envelope glycoprotein G1,	
		cnvelope glycoprotein G2) [Hantavirus sp.=Puumala	
2183	285275	virus, Hallnas strain, Peptide, 1148 aa]	0.6
		,	
		gi 7506378 pir  T23989 hypothetical protein R07A4.3 -	
		Caenorhabditis elegans emb CAA91763.1  (Z67756)	
		cDNA EST yk63e10.5 comes from this gene~cDNA	
2186	7506378	EST yk63c10.3 comes from this gene [Caenorhabditis clegans]	8.9
2100		gi 4511976 gb AAD21536.1  (AF088896) unknown	0.5
2187		[Zymomonas mobilis]	0.65
		gi 14761847 ref XP_017198.2  hypothetical protein	
2188	14761847	FLJ12085 [Homo sapiens]	3E-13
		gi 5835478 ref NP_008404.1 CYTB_13475 cytochrome	
		b [Balanoglossus carnosus] pir  T11138 ubiquinol	
Ì	i	cytochrome-c reductase (EC 1.10.2.2) cytochrome b -	
2100	5025470	acorn worm mitochondrion gb AAD11951.1	0.05
2192	3833478	(AF051097) cytochrome b [Balanoglossus carnosus]	0.95
2195	14006462	gi 14906463 gb AAK72690.1  (AY039648) transcription factor Rel 1 [Crassostrea gigas]	9.3
2193	14900403	gi 6680964 rcf NP 031758.1  procollagen, type XVII,	9.3
		gijoos0964 rer  NP_031738.1  procollagen, type XVII,   alpha 1 [Mus musculus] pir  A46053 bullous	
		pemphigoid antigen, BPAG2, type XVII collagen alpha	
		1-chain - mouse gb AAA37443.1  (L08407) collagen	
2196	6680964	type XVII [Mus musculus]	6.8
-2170	0000701	by po 12 1 11 [11 and museums]	0.0
		gi 7206631 gb AAF39790.1  (AC006631) Hypothetical	
2197	7206631	protein F27B3.2 [Caenorhabditis elegans]	4.3
		gi 2564679 gb AAB81836.1  (AF023484) putative KP78	
2201	2564679	protein kinase [Drosophila melanogaster]	0.83
		gi 9558143 emb CAC00269.1  (AL160371) possible	
2204	9558143	f16d3.1 protein [Leishmania major]	5.3
2205	12012146	gi 13813146 gb AAK40384.1  (AE006643) ATP- dependent helicase [Sulfolobus solfataricus]	4.2
2205	13813146		4.3
		gi 7662168 rcf NP_055497.1  KIAA0535 gene product [Homo sapiens] dbi BAA25461.1  (AB011107)	
2206	7662169	KIAA0535 protein [Homo sapiens]	0.0003
2200	7002100	Exercision from sabiens	0.0003

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 9229890 dbj BAB00618.1  (AB036841) prickle 2	
2214	9229890	[Ciona intestinalis]	4.4
		gi 3435174 gb AAC32342.1  (AF061251) O antigen	
		flippase Wzx [Escherichia coli] gb AAG57097.1 AE005429 8 (AE005429) O antigen	
2217	3435174	flippase Wzx [Escherichia coli O157:H7 EDL933]	2.5
ZZI/	3433174	imppuse WZZ [Escalericina con O15/11/ EDE935]	2.3
		gi 5725923 gb AAD48242,1 AF089987 1 (AF089987)	
		four-loop conotoxin ABVIF [Conus abbreviatus]	
		gb AAD48243.1 AF089988 1 (AF089988) four-loop	
	ĺ	conotoxin ABVIF [Conus abbreviatus]	
		gb AAD48244.1 AF089989 1 (AF089989) four-loop	
		conotoxin ABVIF [Conus abbreviatus]	
	1	gb AAD48245.1 AF089990 1 (AF089990) four-loop	
2225	5725923	conotoxin ABVIF [Conus abbreviatus]	3.8
	1	gi 1705523 sp P52650 C24A_PIG CYTOCHROME B-	
		245 LIGHT CHAIN (P22 PHAGOCYTE B-	
		CYTOCHROME) (NEUTROPHIL CYTOCHROME	
		B, 22 KD POLYPEPTIDE) (P22-PHOX)	
	Į	(CYTOCHROME B(558) ALPHA CHAIN)	
		(SUPEROXIDE-GENERATING NADPH OXIDASE	
2220	1505500	LIGHT CHAIN SUBUNIT) gb AAA64635.1  (U02477)	
2228	1705523	NADPH oxidase light chain subunit [Sus scrofa]	0.48
		gi 730885 sp P07989 T1M_SALPO TYPE I	40
		RESTRICTION ENZYME STYSPI M PROTEIN	
		(M.STYSPI) gb AAA27143.1  (L02507) restriction- modification enzyme type I M subunit [Salmonella	
2240	730885	enterical	4.9
	750005	ontoriouj	7.2
		cil6752573 mgND 0241261  mdochoom P450 24	
		gi 6753572 ref NP_034126.1  cytochrome P450, 24 [Mus musculus] sp Q64441 CP24 MOUSE	
		CYTOCHROME P450-CC24, MITOCHONDRIAL	
		PRECURSOR (P450-CC24) (VITAMIN D(3) 24-	
		HYDROXYLASE) (1,25-DIHYDROXYVITAMIN	
		D(3) 24-HYDROXYLASE) (24-OHASE) pirl S60033	
1		25-hydroxyvitamin D3 24-hydroxylase precursor -	
		mouse dbj[BAA08416.1  (D49438) 25-hydroxyvitamin	
		D3 24-hydroxylase precursor [Mus musculus]	
		dbj BAA21843.1  (D89669) vitamin D-24-hydroxylase	
2241	6753572	[Mus musculus]	9.7

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 14578299 gb AAF99465.1  (AY003872)	
2243	14578299	PV1H14095_P [Plasmodium vivax]	3.7
		gi 1916617 gb AAB51194.1  (U66003) ADAM 13	
2245	1916617	[Xenopus laevis]	0.45
		gi 5739073 gb AAD50327.1  (AF063693) type XIII	
2249	5739073	collagen [Mus musculus]	2.5
l		gi 3135611 gb AAC29067.1  (AF062485) cellulose	
2254	3135611	synthase [Arabidopsis thaliana]	3.1
		gi 281689 pir  S27657 hypothetical protein 1 -	
		Rhizobium meliloti gb AAA26255.1  (M94085) not	
		homologous to known sequences as of 2/92; ORF1;	
2256	281689	putative [Sinorhizobium meliloti]	6
		gi 10434352 dbj BAB14232.1  (AK022759) unnamed	
2257	10434352	protein product [Homo sapiens]	2E-23
		gi 1504022 dbj BAA13210.1  (D86974) KIAA0220	_
2258	1504022	[Homo sapiens]	4E-21
		gi]1871176 gb AAB63536.1  (U90439) unknown protein	
2266	1871176		7.5
		gi 12846015 dbj BAB26996.1  (AK010513) putative	
2270	12846015	[Mus musculus]	8
		gi 12860337 dbj BAB31923.1  (AK019929) putative	
2273	12860337	[Mus musculus]	5
1	1		
		gi 994736 gb AAA75561.1  (M18327) LacOPZ-alpha	
		peptide from pUC9; putative [unidentified cloning	
		vector] gb AAA75563.1  (M18328) LacOPZ-alpha	
		peptide from pUC9; putative [Cloning vector pBGS9+]	
0076	004506	gb AAA75565.1  (M18329) LacOPZ-alpha peptide from	
2276	994736	pUC9; putative [Cloning vector pBGS9-]	0.00002
		gi 11352313 pir  G83376 probable trehalose synthase	
		PA2152 [imported] - Pseudomonas aeruginosa (strain	
2280	11252212	PAO1) gb AAG05540.1 AE004642_7 (AE004642) probable trehalose synthase [Pseudomonas aeruginosa]	
2200	11332313	probable trendiose symmase [Pseudomonas aeruginosa]	2
		ail7494002laidfF00921 kymathati1i 75127 14 5	
		gi 7486992 pir  T00831 hypothetical protein T13L16.5 - Arabidopsis thaliana gb AAD20114.2  (AC006201)	
2285	7486992	hypothetical protein [Arabidopsis thaliana]	1.4
2203	7-100332	gi 10173203 dbj BAB04308,1  (AP001509)	1.4
		BH0589~unknown conserved protein in others [Bacillus	
2286	10173203	halodurans]	7.3
			1.5

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
	ł		
		gi 7497857 pir  T20180 hypothetical protein C53B4.4a -	
l		Caenorhabditis elegans emb CAA92457.1  (Z68215)	
		contains similarity to Pfam domain; PF00130 (Phorbol	
		esters/diacylglycerol binding domain (C1 domain)),	
	ŀ	Score=12.8, E-value=0.0015, N=1; PF00595 (PDZ	
		domain (Also known as DHR or GLGF).), Score=34.8,	
2289	7497857		9.6
		gi 6424831 gb AAF08166.1  (AF130210) NADH	
2291	6424831	dehydrogenase subunit F [Impatiens biflora]	6.1
		gi 7489002 pir  T07021 extensin-like protein Dif10	
		precursor - tomato (fragment) emb CAA67813.1	
		(X99451) extensin-like protein Dif10 [Lycopersicon	
2292	7489002		5.6
		gi 1684828 gb AAB36537.1  (U77681) tyrosine kinase	
2294	1684828	receptor [Xenopus laevis]	2
		gi 12859724 dbj BAB31753.1  (AK019486) putative	
2297	12859724	[Mus musculus]	0.0003
		gi 11498284 ref[NP_069510.1  adenylate kinase (adk)	
		[Archaeoglobus fulgidus] sp O29581 KAD_ARCFU	
		ADENYLATE KINASE (ATP-AMP	
		TRANSPHOSPHORYLASE) pir  D69334 adenylate	
		kinase (EC 2.7.4.3) - Archaeoglobus fulgidus	
		gb AAB90565.1  (AE001058) adenylate kinase (adk)	
2301	11498284	[Archaeoglobus fulgidus]	3E-13
		gi 14725330 ref XP_002254.2  mitochondrial	
		translational initiation factor 2 precursor [Homo	
2309	14725330		4.5
		gi 79651 pir  A30189 iron stress-induced hypothetical	
2313	79651	protein precursor - Synechococcus sp	2.8
		,,	
		gi 7500007 pir  T16186 hypothetical protein F27D9.4 -	
	7500007	Caenorhabditis elegans gb AAA93383.1  (U49829)	
2314	7500007	Hypothetical protein F27D9.4 [Cacnorhabditis elegans]	7.1
2217	12517022	gi 13517833 gb AAK29011.1  (AF344620) long-	
2317	13517833	wavelength rhodopsin [Ceratina calcarata]	3.4

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
	ľ	gi 7478661 pir  F70662 probable plcC protein -	
ļ		Mycobacterium tuberculosis (strain H37RV)	
		emb CAB06146.1  (Z83860) plcC [Mycobacterium	
		tuberculosis] gb AAK46707.1  (AE007081)	
		phospholipase C [Mycobacterium tuberculosis	
2318	7478661	CDC1551]	4.4
		gi 14768227 ref XP_012121.3  purinergic receptor P2X,	
2323	14768227	ligand-gated ion channel, 7 [Homo sapiens]	4.3
		gi 283032 pir  S22456 hydroxyproline-rich glycoprotein -	
		perennial teosinte emb CAA45514.1  (X64173)	
2328	283032	hydroxyproline-rich glycoprotein [Zea diploperennis]	5.4
		gi 10241645 emb CAC09484.1  (AL442113) putative	
2329	10241645	protein [Oryza sativa]	9.1
		gi 14733085 ref XP_003575.3  soluble liver antigen/liver	
2330	14733085	pancreas antigen [Homo sapiens]	1E-17
		gi 12188796 emb CAC21494.1  (AJ278866) MchF	
2335	12188796	protein [Escherichia coli]	4.7
		gi 2492604 sp P78595 CDR2_CANAL MULTIDRUG	
		RESISTANCE PROTEIN CDR2 gb AAB96797.1	
2340	2492604	(U63812) drug resistance protein 2 [Candida albicans]	6.4
		gi 13365569 dbj BAB39114.1  (AP002897) hypothetical	
		protein~similar to Oryza sativa chromosome 1,	
2345		P0665D10.16	1.8
		gi 1334612 emb CAA41034.1  (X57968) nad1 [Triticum	
2346	1334612		4.8
		gi 1334612 cmb CAA41034.1  (X57968) nad1 [Triticum	
2351	1334612		4.8
	100	gi 1334612 emb CAA41034.1  (X57968) nad1 [Triticum	
2352	1334612		4.6
2252	1204612	gi 1334612 emb CAA41034.1  (X57968) nad1 [Triticum	
2353	1334612		4.8
2256	1004610	gi 1334612 cmb CAA41034.1  (X57968) nad1 [Triticum	
2356	1334612		4.4
		gi 14250644 gb AAH08786.1 AAH08786 (BC008786)	
2357	14250744	integrin, alpha 5 (fibronectin receptor, alpha	
2337	14230644	polypeptide) [Homo sapiens]	2
2367	7202054	gi 7293054 gb AAF48440.1  (AE003498) Top1 gene	
2307	1293054	product [Drosophila melanogaster]	1.2
2369	13050244	gi 13959344 sp P82957 DM43_DIDMA VENOM	
2309	13939344	METALLOPROTEINASE INHIBITOR DM43	8.2

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		Topped and the transfer of the	
2271	0000017	gi 9082017 gb AAF82689.1 AF227196_1 (AF227196)	
2371	9082017	ORF1a polyprotein [gill-associated virus]	7.8
	l		
		gi 7498863 pir  T20730 hypothetical protein F10G8.8 -	
		Caenorhabditis elegans emb CAB02286.2  (Z80216)	
		Weak similarity with intermediate filament protein	
		(TREMBL id G633240), contains similarity to Pfam	
		domain: PF00169 (PH domain), Score=79.3, E-	
		value=2.7e-21, N=2~cDNA EST EMBL:T01262 comes	
		from this gene~cDNA EST yk23d5.3 comes from this gene~> emb CAA19441.2  (AL023823) Weak similarity	
		with intermediate filament protein (TREMBL id	
		G633240), contains similarity to Pfam domain:	
		PF00169 (PH domain), Score=79.3, E-value=2.7e-21,	
		N=2~cDNA EST EMBL:T01262 comes from this	
2374	7498863	gene~cDNA EST yk23d5.3 comes from this gen>	2.5
		gi 321514 pir  S27931 Env/v-mpl fusion protein -	
		myeloproliferative leukemia virus gblAAA77654.1	
		(M60350) env:v-mpl fusion protein [Mycloproliferative	
2375	321514	leukemia virus]	2.8
		gi 11348760 pir  C83635 hypothetical protein PA0086	
		[imported] - Pseudomonas aeruginosa (strain PAO1)	
		gb AAG03476.1 AE004447_9 (AE004447) hypothetical	
2376	11348760	protein [Pseudomonas aeruginosa]	4.6
		gi 742234 prf  2009317A adhA upstream ORF	
2377	742234	[Acctobacter (subgen. Acctobacter) aceti]	8.2
		gi 11466189 ref NP_066512.1  apocytochrome b	
2379	11466100	[Naegleria gruberi] gb AAG17790.1 AF288092_15	7.0
2319	11400189	(AF288092) apocytochrome b [Naegleria gruberi] gi 8250181 cmb CAB93524.1  (AJ271740) D-Titin	7.2
2380	8250181	[Bulgas 25018   Cambi Cab 93524.1   (AJZ/1740) D-11tin  [Drosophila melanogaster]	4,2
2380	0230101	gi 5870848 gb AAB03857.2  (U31864) stearyl-CoA	4,2
2383	5870848	desaturase [Cyprinus carpio]	7.7
2505	3070040	gi 4063751 gb AAC98459.1  (AC005851) putative ABC	7.7
2387	4063751	transporter [Arabidopsis thaliana]	7.6
		gi 7504822 pir  T33371 hypothetical protein H02F09.1 -	
		Caenorhabditis elegans gb AAC64621.1  (AF077538)	
2389	7504822	unknown [Caenorhabditis clegans]	3,6

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
NO	-11	gi 15011889 ref[NP 077251.1  oxysterol binding protein	P VALUE
		2 [Mus musculus] emb CAC16404.2  (AJ278263)	
		oxystyrol-binding protein homologue 1 [Mus musculus	
2390	15011889		1.9
2390	13011889		1.9
		gi 450730 cmb CAA50838.1  (X71982) ORF j18L; potential membrane spanning region; potential	
2393	450730	glcosylation site [African swine fever virus]	2.5
2393	430730		2.3
		gi 7445990 pir  G72290 branched chain amino acid ABC	
		transporter, ATP-binding protein - Thermotoga	
		maritima (strain MSB8) gb AAD36215.1 AE001771_8	
		(AE001771) branched chain amino acid ABC	
2200	7445000	transporter, ATP-binding protein [Thermotoga	
2398	7445990		7.4
		gi 11990448 dbj BAB19782.1  (AB052747) vascular	
		cell adhesion molecule-1 6D variant lacking D7 [Bos	
2401	11990448		4.6
	l	gi 12841678 dbj BAB25308.1  (AK007856) putative	
2402	12841678	[Mus musculus]	1E-39
	<b>.</b>	gi 5052967 gb AAD38786.1 AF151533_1 (AF151533)	
2403	5052967	polyketide synthase [Nodulisporium sp. ATCC74245]	6.3
		gi 8099350 gb AAF72105.1 AF154847_1 (AF154847)	
2407	8099330	33 kDa Vamp-associated protein [Homo sapiens]	5.2
		gi 267344 sp P29791 VGLF_BRSVA FUSION	
		GLYCOPROTEIN PRECURSOR [CONTAINS:	
		FUSION GLYCOPROTEIN F2; FUSION	
		GLYCOPROTEIN F1] pir  VGNZBA cell fusion	
		glycoprotein precursor - bovine respiratory syncytial	
		virus (strain A51908) gb AAA42804.1  (M82816)	
2410	267344	fusion protein F [Bovine respiratory syncytial virus]	1.4
		gi 12856615 dbj BAB30727.1  (AK017396) putative	
2417	12856615		8E-49
	1	gi 11347010 pir  B81303 probable membrane protein	
		Cj1013c [imported] - Campylobacter jejuni (strain	
0.405		NCTC 11168) emb CAB73269.1  (AL139077) putative	
2425	11347010	membrane protein [Campylobacter jejuni]	0.89
	İ		2
		gi 7499991 pir T29526 hypothetical protein F27C1.7 -	
	1	Caenorhabditis elegans gb AAB37654.1  (U80441)	
2426	7499991	Hypothetical protein F27C1.7 [Caenorhabditis elegans]	3.1

	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS			
NO	N	DESCRIPTION	P VALUE	
1		gi 2498043 sp Q57568 Y104_METJA		
		HYPOTHETICAL ATP-BINDING PROTEIN MJ0104		
1		pir H64312 probable DNA helicase MJ0104 -		
		Methanococcus jannaschii gb AAB98084.1  (U67467)		
2427	0.4000.40	DNA-binding protein, probably DNA helicase		
2421	2498043		1.5	
		gi 7494263 pir  T18488 hypothetical protein C0825c -		
ĺ		malaria parasite (Plasmodium falciparum)		
		cmb CAB11127.1  (Z98551) putative cleavage and polyadenylation specificity factor protein [Plasmodium		
2430	7/10/263	falciparum]	2.3	
2430	7494203	gi 493224 dbi BAA03434.1  (D14581) fatty-acid	2.3	
2434	493224	desaturase [Anabaena variabilis]	9.1	
2434	473224	ucsaturase [miabaciia variabilis]	9.1	
		gi 6755468 ref NP 036019.1  septin 3 [Mus musculus]		
		spiQ9Z1S5 SEP3 MOUSE NEURONAL-SPECIFIC		
		SEPTIN 3 gb AAD02884.1  (AF104411) neuronal-		
2438	6755468	specific septin 3 [Mus musculus]	0.59	
		gi 9967295 dbj BAB12347.1  (AB047936) hypothetical	5.55	
2440	9967295	protein [Macaca fascicularis]	0.063	
		gi 12847975 dbj BAB27780.1  (AK011690) putative		
2447	12847975	[Mus musculus]	7E-65	
		gi 1730946 sp P50833 YPPE BACSU		
		HYPOTHETICAL 14.5 KDA PROTEIN IN PONA-		
		COTD INTERGENIC REGION pir  B69940		
		hypothetical protein yppE - Bacillus subtilis		
		gb AAB38463.1  (L47838) putative [Bacillus subtilis]		
2451	1730946	emb CAB14144.1  (Z99115) yppE [Bacillus subtilis]	9.3	
		gi 401192 sp P30975 TLR2_DROME TACHYKININ-		
i I		LIKE PEPTIDES RECEPTOR 99D (DTKR)		
		pir  S17783 tachykinin receptor homolog DTKR - fruit		
		fly (Drosophila melanogaster) emb CAA44595.1		
		(X62711) receptor for tachykinin-like peptides		
2452	401192	[Drosophila mclanogaster]	4.8	
2452		gi 14768202 ref XP_018137.2  L1 cell adhesion	0.11	
2453	14/68202	molecule precursor [Homo sapiens]	0.11	

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE	
2457	141028	gi 141028 sp P04540 NUSM_TRYBB NADH- UBIQUINONE OXIDOREDUCTASE CHAIN 5 pir  QQUTC5 NADH dehydrogenase (ubiquinone) (EC 1.6.5.3) chain 5 - Trypanosoma brucei mitochondrion gb AAB59225.1  (M14820) NADH dehydrogenase subunit 5 [Trypanosoma brucei] emb CAB57807.1  (X01094) unidentified reading frame 10 [Trypanosoma brucei]	3.6	
		gi 7499039 pir  T20867 hypothetical protein F13H10.5 - Caenorhabditis elegans emb CAA92956.1  (Z68748) contains similarity to Pfam domain: PF01663 (Type I phosphodiesterase / nucleotide pyrophosphatase), Score-512.3, E-value-1.1e-150, N=1 [Caenorhabditis elegans] emb CAA15977.1  (ALD21176) contains similarity to Pfam domain: PF01663 (Type I phosphodiesterase / nucleotide pyrophosphatase), Score-512.3, E-value-1.1e-150, N=1 [Caenorhabditis		
2461	7499039	elegans]	9.7	
2462	3023956	gij3023956 sp Q00808 HET1_PODAN VEGETATIBLE INCOMPATIBILITY PROTEIN HET-E-1 pir  T18521 beta transducin-like protein - Podospora anserina gb AAA85775.1  (L28125) beta transducin-like protein [Podospora anserina]	8.7	
2463	14972564	gi 14972564 gb AAK75201.1  (AE007410) glutamine amidotransferase, class I [Streptococcus pneumoniae]	0.4	
2470	13928966	gij 13928966 ref NP_113882.1  heat shock factor 2 [Rattus norvegicus] gbjAAD51329.1 AF172640_1 (AF172640) heat shock factor 2 [Rattus norvegicus] gij4521320dbjiBAA11580.1  (082816) product is	6E-14	
2473	4521320	gi 4521320 dbj BAA11580.1  (D82816) product is unknown [Gallus gallus]	0.29	
2475	7507534	gij7507534[pii][T24738 hypothetical protein T09E11.2 - Caenorhabditis elegans emb](CAB03532.1] (Z81147) Similarity to zine finger proteins, contains similarity to Pfam domain: PF00104 (Ligand-binding domain of nuclear hormone receptor), Score=14.1, E- value=0.00047, N=1; PF00105 (Zine finger, C4 type (two domains)), Score=42.6, E-value=5.5e-12, N>	6.7	

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS			
NO	N	DESCRIPTION	P VALUE	
		gi 927030 gb AAA73871.1  (L13287) kcrB4 gene		
2478	927030	product [Escherichia coli]	3,5	
		gi 6724309 gb AAF26929.1  (AF079967) NADH		
2481	6724309	dehydrogenase subunit 4 [Phytomonas serpens]	7.9	
		l i		
		gi 6319700 rcf NP_009783.1  Ybr224wp		
	ļ	[Saccharomyces cerevisiae] sp P38320 YB74_YEAST		
		HYPOTHETICAL 19.3 KDA PROTEIN IN FAT2-		
		MCX1 INTERGENIC REGION PRECURSOR		
		pir  S46100 probable membrane protein YBR224w -		
		yeast (Saccharomyces cerevisiae) emb CAA85187.1		
2482	6319700	(Z36092) ORF YBR224w [Saccharomyces cerevisiae]	2.4	
		gi 7503173 pir  T31884 hypothetical protein F41E6.14 -		
		Caenorhabditis elegans gb AAB65962.1  (AF016448)		
		weak similarity to several acyltransferases		
2496	7503173	[Caenorhabditis elegans]	9.7	
		gi 11352621 pir  E83187 tetrahydrodipicolinate		
		succinylase PA3666 [imported] - Pseudomonas		
	1	aeruginosa (strain PAO1) dbj BAA75911.1		
		(AB024601) tetrahydrodipicolinate N-		
		succinyletransferase [Pseudomonas aeruginosa]		
		gb AAG07054.1 AE004786_6 (AE004786)		
		tetrahydrodipicolinate succinylase [Pseudomonas		
2498	11352621	aeruginosa]	3.3	
		gi 7662214 ref NP_055604.1  KIAA0628 gene product		
		[Homo sapiens] ref XP_005044.3  KIAA0628 gene		
		product [Homo sapiens] dbj BAA31603.1  (AB014528)		
2500	7662214	KIAA0628 protein [Homo sapiens]	3	
		gi 7508531 pir  T25325 hypothetical protein T26H2.7 -		
		Caenorhabditis elegans emb CAB04848.1  (Z82055)		
		contains similarity to Pfam domain: PF01757 (Domain		
		of unknown function), Score=543.5, E-value=4.6e-160,		
2506	7508531	N=1 [Caenorhabditis elegans]	9.2	
		gi 14730527 ref XP_051896.1  phospholipase A2, group		
		IVA (cytosolic, calcium-dependent) [Homo sapiens]		
2505		ref[XP_051897.1] phospholipase A2, group IVA		
2507	14730527	(cytosolic, calcium-dependent) [Homo sapiens]	5E-26	

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#### Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins) ACCESS SEO ID N NO DESCRIPTION P VALUE gi|138394|sp|P27330|VHEL_LSV PROBABLE HELICASE (ORF 2) emb|CAA33398.1| (X15343) 2510 138394 25kD protein [Lily symptomless virus] 10 gi|13938116|gb|AAH07173.1|AAH07173 (BC007173) Unknown (protein for IMAGE:3493127) [Mus 2519 13938116 musculus1 8E-22 gi|7486371|pir|T00661 hypothetical protein F3I6.24 -Arabidopsis thaliana gb|AAC00591.1| (AC002396) 2520 7486371 Unknown protein [Arabidopsis thaliana] 0.16 gil11358141|pir||T48198 hypothetical protein T20L15.40 - Arabidopsis thaliana emblCAB82747.11 2523 11358141 (AL162351) putative protein [Arabidopsis thaliana] 9.1 gil14485227lgblAAK62977.1lAF384372 3 (AF384372) 2526 14485227 surface antigen [Hepatitis B virus] 2.6 gil6594617|gb|AAF18559.1|U42380 1 (U42380) 6594617 aminopeptidase [Aplysia californica] 2527 5.3 gi|14193306|gb|AAK55890.1|AF267211 2 (AF267211) ATP synthase gamma subunit [Candidatus Carsonella 2529 14193306 ruddiil 3.5 gil3319680|emb|CAA76809.1| (Y17614) N8 protein 2531 3319680 [Medicago truncatula] 1.5 gi|13811987|ref[NP 113116.1| DNA renair helicase component of transcription factor b [Guillardia theta] gb|AAK39689.1|AF083031_46 (AF083031) DNA repair helicase component of transcription factor b 2536 13811987 [Guillardia theta] 4.8 gi|3122601|sp|P93107|PF20 CHLRE FLAGELLAR WD-REPEAT PROTEIN PF20 pir T08180 PF20 protein, microtubule-associated - Chlamydomonas reinhardtii gb|AAB41727.1| (U78547) PF20 2537 3122601 [Chlamydomonas reinhardtii] 1.1 gil7460001|pirl|G71079 hypothetical protein PH0903 -Pyrococcus horikoshii dbi|BAA29997.1| (AP000004) 2540 7460001 141aa long hypothetical protein [Pyrococcus horikoshii] 6.3 gi|12330702|gb|AAG52889,1|AF333769 1 (AF333769) 2541 12330702 cell recognition molecule CASPR3 [Homo sapiens] 2E-28

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteing)
SEQ ID	ACCESS	Table 5D Treatest freignoof (Blasca vs. 1401-Redundant	r Iotenis)
NO	N	DESCRIPTION	P VALUE
		gi 6754652 ref NP_034904.1  methyl-CpG binding	
		domain protein 4 [Mus musculus] gb[AAC68878.1]	
		(AF072249) methyl-CpG binding protein MBD4 [Mus	
		musculus gb AAD56595.1 AF120996 1 (AF120996)	
1		methyl-CpG binding protein 4 [Mus musculus	
2547	6754652	domesticus]	7.1
		gi 12655370 emb CAB57344.3  (AJ243708) prickle pk	
2548	12655370	isoform [Drosophila melanogaster]	7.7
		gi 12853631 dbj BAB29800.1  (AK015333) putative	
2551	12853631		0.61
		gi 11466216 ref[NP_066539.1  hacm lyase [Naegleria	
		gruberi] gb AAG17817.1 AF288092_42 (AF288092)	
2554	11466216	haem lyase [Naegleria gruberi]	1.5
		gi 7321597 gb AAA32099.2  (L28677) unknown	
2557	7321597	[Tetrahymena pyriformis]	3
		gi 7304202 gb AAF59238.1  (AE003840) CG1602 gene	
2560	7304202	product [Drosophila melanogaster]	1.6
		gi 336831 gb AAB02281.1  (M57910) NADH	
2564	336831	dehydrogenase subunit 2 [Drosophila melanogaster]	5.5
		gi 7515231 pir  T13518 hypothetical protein 29 -	
		Bacillus phage phi-105 dbj BAA36635.1  (AB016282)	
2566	7515231		7.7
25.00	10045043	gi 12847263 dbj BAB27500.1  (AK011258) putative	
2568	12847263	[Mus musculus]	6,6
		gi 9964395 ref NP_064863.1  AMV081 [Amsacta	
		moorei entomopoxvirus] gb AAG02787.1 AF250284_81	
2569	9964395	(AF250284) AMV081 [Amsacta moorei entomopoxvirus]	2.4
2309	9904393	gi 12060849 gb AAG48266.1 AF308299 1 (AF308299)	2.4
		serologically defined breast cancer antigen NY-BR-85	
2573	12060849		0,0006
2313	12000047	gi 9633381 ref NP 050485.1  D5L protein [variola	0,000
		minor virus] pir  H72173 D5L protein - variola minor	
		virus (strain Garcia-1966) gb[AAA69395.1] (U18339)	
		D4L [Variola virus] emb[CAA50966.1] (X72086)	
		ORF15L; B16L in citation [3] [Variola virus]	
		emb CAB54786.1  (Y16780) D5L protein [variola	
2575	9633381	minor virus]	7.5
		gi 7407131 gb AAF61923.1  (AF228524) SanE	
2576	7407131	[Streptomyces ansochromogenes]	0.53
		gi 14760789 ref XP_044332.1  piwi (Drosophila)-like 1	
2577	14760789	[Homo sapiens]	9E-42

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 2764800 emb CAA54153.1  (X76738) 12s globulin	
2584	2764800	[Avena sativa]	1.4
		gi 12860337 dbj BAB31923.1  (AK019929) putative	
2585	12860337	[Mus musculus]	5.3
		gi 7768484 emb CAB90775.1  (AL355632) putative	
		mitochondrial carrier protein [Schizosaccharomyces	
2586	7768484	pombe]	2.8
		gi 12620098 gb AAG60558.1  (AF250768) BioA-like	
2587	12620098	protein [uncultured bacterium pCosFS1]	8.4
		gi 2281181 gb AAB66275.1  (U58587) maturase	
2591	2281181	[Lagoecia cuminoides]	3.8
		gi 6573736 gb AAF17656.1 AC009398_5 (AC009398)	
2595	6573736	F20B24.10 [Arabidopsis thaliana]	6.7
		gi 31155 emb CAA24999.1  (X00176) preproenkephalin	
2596	31155	part 1 [Homo sapiens]	9.5
		gi 11278033 pir  C81832 transferrin-binding protein A	
		NMA2024 [imported] - Neisseria meningitidis (group A	
		strain Z2491) gb AAC13726.1  (AF058689) transferrin	
		binding protein A precursor [Neisseria meningitidis]	
		emb CAB85243.1  (AL162757) transferrin-binding	
2600	11278033	protein A [Neisseria meningitidis Z2491]	5
		gi 7493138 pir  T37964 probable ubiquitin ligase -	
		fission yeast (Schizosaccharomyces pombe)	
		emb CAB16714.1  (Z99531) putative ubiquitin ligase	
2613	7493138	[Schizosaccharomyces pombe]	0.21
	*	gi 14625275 gb AAA80360.2  (U39644) Hypothetical	
2615	14625275	protein T10E10.4 [Caenorhabditis elegans]	5.3
		gi 7515479 pir  S72298 hypothetical protein 91 -	
		Plasmodium falciparum plastid emb CAA64588.1	
2617	7515479	(X95276) ORF91 [Plasmodium falciparum]	0.66
		gi 14091855 gb AAK53858.1 AC016781_12	
2619	14091855	(AC016781) Hypothetical protein [Oryza sativa]	0.69
		gi 8393165 ref NP_035661.2  transiently-expressed	
2621	8393165	axonal glycoprotein [Mus musculus]	0.11
		gi 6754242 ref NP_034603.1  histidine rich calcium	
		binding protein [Mus musculus]	
		gb AAD55250.1 AF158597_1 (AF158597) histidine-	
2624	6754242	rich Ca2+ binding protein [Mus musculus]	8.1
		gi 7959261 dbj BAA96024.1  (AB040933) KIAA1500	
2625	7959261	protein [Homo sapiens]	1E-36

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 5508828 gb AAD43995.1  (U59485) AttU	
2626	5508828	[Agrobacterium tumefaciens]	0.27
	ļ	gi 3643603 gb AAC42250.1  (AC005395) unknown	
2627 -	3643603	protein [Arabidopsis thaliana]	4.7
1			
		gi 5453906 ref NP 006310.1  CDP-diacylglycerol	
}		inositol 3-phosphatidyltransferase (phosphatidylinositol	
1		synthase) [Homo sapiens] ref[XP 008065,1] CDP-	
		diacylglycerol-inositol 3-phosphatidyltransferase	
		(phosphatidylinositol synthase) [Homo sapiens]	
1		ref[XP 043951.1] CDP-diacylglycerolinositol 3-	
		phosphatidyltransferase (phosphatidylinositol synthase)	
		[Homo sapiens] sp O14735 PIS HUMAN CDP-	
		DIACYLGLYCEROL-INOSITOL 3-	
		PHOSPHATIDYLTRANSFERASE	
		(PHOSPHATIDYLINOSITOL SYNTHASE) (PTDINS	
		SYNTHASE) (PI SYNTHASE) gb AAB94860.1	
		(AF014807) phosphatidylinositol synthase [Homo	
		sapiens] gb AAH01444.1 AAH01444 (BC001444) CDP-	
		diacylglycerolinositol 3-phosphatidyltransferase	
2630	5453906	(phosphatidylinositol synthase) [Homo sapiens]	7
		gi 7515479 pir  S72298 hypothetical protein 91 -	
		Plasmodium falciparum plastid emb CAA64588.1	
2631	7515479	(X95276) ORF91 [Plasmodium falciparum]	0.71
		gi 11467075 ref NP_042551.1  ribosomal protein L5	
		[Acanthamoeba castellanii] sp P46764 RM05_ACACA	
		MITOCHONDRIAL 60S RIBOSOMAL PROTEIN L5	
		pir  S53852 ribosomal protein L5 - Acanthamoeba	
		castellanii mitochondrion gb AAD11844.1  (U12386)	
2636	11467075	ribosomal protein L5 [Acanthamoeba castellanii]	2.3
		gi 1352549 sp P48906 NU2M_HANWI NADH-	
2639	1352549	UBIQUINONE OXIDOREDUCTASE CHAIN 2	4.1
		gi 13111580 gb AAK12385.1 AF296091_1 (AF296091)	
2642	13111580	polyprotein [Porcine teschovirus]	2.4
		gi 1314734 gb AAA99804.1  (U54641) 220 kDa silk	
2646	1314734	protein [Chironomus thummi]	1.9
		gi 7001374 gb AAF34871.1 AF112184_1 (AF112184)	
		serine/threonine kinase NKIATRE alpha [Rattus	
2651	7001374	norvegicus]	2.9
		gi 2687582 gb AAB88853.1  (AF032875) protein kinase	
2653	2687582	[Mus musculus]	7.2

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 10636263 emb CAC10528.1  (AJ293919) putative	
0.654		inositol 1,4,5-trisphosphate receptor [Caenorhabditis	_
2654	10636263	briggsac	5
		gi 5902891 dbj BAA84474.1  (AB032367) type I	
2663	5902891	polyketide synthase AVES 1 [Streptomyces avermitilis]	0.087
2003	3702071	gi 4567214 gb AAD23629.1 AC007113 2 (AC007113)	0.087
2664	4567214		6.4
2001	1507211	gi 2326816 emb CAA99382.1  (Z75081) ORF	0.4
2665	2326816	YOR172w [Saccharomyces cerevisiae]	0.46
		, , , , , , , , , , , , , , , , , , , ,	
		gi 14285987 sp O83933 Y967 TREPA	
		HYPOTHETICAL PROTEIN TP0967 pirlB71260	
		hypothetical protein TP0967 - syphilis spirochete	
		gb AAC65925.1  (AE001264) T. pallidum predicted	
2667	14285987		6.9
		gi 6678247 ref[NP 033358.1  transcription factor 7-like	
		1 [Mus musculus] emb CAA11070.1  (AJ223069) TCF-	
2669	6678247	3 protein [Mus musculus]	3.4
		gi 7491697 pir  T40527 hypothetical protein	
		SPBC530.11c - fission yeast (Schizosaccharomyces	
		pombe) emb CAA19177.1  (AL023634) putative	
2671	7491697	transcriptional regulator [Schizosaccharomyces pombe]	4.9
0.000		gi 5459308 emb CAB50693.1  (AJ238951) CE9 protein	
2673	5459308	[Canis familiaris]	8.8
2675	7959261	gi 7959261 dbj BAA96024.1  (AB040933) KIAA1500	
2673	7939261	protein [Homo sapiens]	1E-36
		gi 14520279 ref NP_125754.1  hypothetical protein	
		[Pyrococcus abyssi] pir  B75192 hypothetical protein	
		PAB2304 - Pyrococcus abyssi (strain Orsay) cmb CAB48985.1  (AJ248283) hypothetical protein	
2680	14520270	[Pyrococcus abyssi]	6.7
2000	14320219	gi 12654531 gb AAH01098,1 AAH01098 (BC001098)	0.7
		Unknown (protein for IMAGE:3508043) [Homo	
2681	12654531	sapiens]	2.8
		gi 12657687 gb AAK01000,1  (AF178873) NADH	2.0
2682	12657687	dehydrogenase subunit 1 [Archiearis parthenias]	0.68

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
	1		
	1	gi 6319517 ref NP_009599.1  Ybr043cp	
		[Saccharomyces cerevisiae] sp P38227 YBP3_YEAST	
		HYPOTHETICAL 77.3 KDA PROTEIN IN FAT1-	
		TCM62 INTERGENIC REGION pir S45901 probable	
	l	membrane protein YBR043c - yeast (Saccharomyces	
		cerevisiae) emb CAA84985.1  (Z35912) ORF YBR043c	
2686	6319517	[Saccharomyces cerevisiae]	7.2
		gi 7019565 ref NP_037528.1  ubiquitin specific protease	
1		25; ubiquitin specific protease USP25 [Homo sapiens]	
1	-	ref XP_047738.1  ubiquitin specific protease 25 [Homo	
		sapiens] gb AAF24998.1 AF134213_1 (AF134213)	
2691	7019565	ubiquitin-specific protease [Homo sapiens]	2E-20
		gi 2130214 pir  S67381 tubulin-folding cofactor D	1
		homolog - fission yeast (Schizosaccharomyces pombe)	
		pir  T39319 tubulin-folding cofactor D homolog - fission	
		yeast (Schizosaccharomyces pombe) emb CAA20686.1	
		(AL031528) tubulin-folding cofactor d.	
2693	2130214	[Schizosaccharomyces pombe]	3.9
		gi 2120601 pir  JC6030 3-oxo-5alpha-steroid 4-	
		dehydrogenase (EC 1.3.99.5) - Comamonas testosteroni	
2696	2120601	gb AAB08517.1  (L23428) delta 4, 5-alpha steroid	146
2696	2120601	dehydrogenase [Comamonas testosteroni]	4.6
		-140/05041-114 A D15050 11 (4/000404011	
1		gi 4263524 gb AAD15350.1  (AC004044) hypothetical protein [Arabidopsis thaliana] emb CAB77761.1	
2697	4263524	(AL161495) hypothetical protein [Arabidopsis thaliana]	5.8
2031	7203324	gi 5453072 gb AAD43426.1  (AF073977) olfactory	5.0
2698	5453072	receptor [Mus musculus domesticus]	1.1
2070	2 133072	gi 7298915 gb AAF54120.1  (AE003675) CG10267	1.1
2701	7298915	gene product [Drosophila melanogaster]	0.86
		gi 158148 gb AAA28827.1  (M19537) RNA polymerase	
		II largest subunit (, EC 2.7.7.6) [Drosophila	
2703	158148	melanogasterl	0.95
		gi 7492334 pir  T37965 probable 40s ribosomal protein -	
		fission yeast (Schizosaccharomyces pombe)	
		emb CAB16715.1  (Z99531) putative component of U3	
[ i		snoRNP, required for pre-18S rRNA processing	
2706	7492334	[Schizosaccharomyces pombe]	6.8
2706	7492334		6.8

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#### Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins) SEO ID ACCESS NO Ν DESCRIPTION P VALUE gil9366627[emblCAB95389.1] (AL359782) conserved 2709 9366627 hypothetical protein, CHR1.84 [Trypanosoma brucei] 9.2 gi|7436828|pir|T00129 hypothetical protein 7 -Leptospira interrogans (fragment) dbj[BAA24376.1] 2713 7436828 (AB010203) ORF7; putative [Leptospira interrogans] 7.2 gi|3805770|gb|AAC69148.1| (U78721) auxin response transcription factor 3 (ETTIN/ARF3) [Arabidopsis thaliana] gb|AAG53998.1|AF336917 1 (AF336917) ARF3 [Arabidopsis thaliana] gb|AAK26023.1|AF360313 1 (AF360313) ARF3 2715 3805770 [Arabidopsis thaliana] 0.65 gil5454158|reflNP 006286.1| valvl-tRNA synthetase 2 [Homo sapiens] splP26640|SYV2 HUMAN VALYL-TRNA SYNTHETASE 2 (VALINE-TRNA LIGASE 2) (VALRS 2) gb|AAD21819.1| (AF134726) G7A 2716 5454158 [Homo sapiens] 1.6 gi|7019565|ref|NP 037528.1| ubiquitin specific protease 25; ubiquitin specific protease USP25 [Homo sapiens] ref[XP 047738.1] ubiquitin specific protease 25 [Homo sapiens] gb|AAF24998.1|AF134213 1 (AF134213) 2721 7019565 ubiquitin-specific protease [Homo sapiens] 8E-21 gi|13186335|gb|AAK15378.1| (AF211133) tryptophanyl-2722 13186335 tRNA synthetase [Carsonella ruddii] 8.9 gil7514607|pir||A71301 conserved hypothetical protein TP0636 - syphilis spirochete gblAAC65610.11 (AE001238) conserved hypothetical protein [Treponema 2726 7514607 pallidum] 9.4 gi|13129046|ref|NP 076972.1| hypothetical protein MGC2718 [Homo sapiens] ref[XP 017742.2] hypothetical protein MGC2718 [Homo sapiens] ref[XP 043040.1| hypothetical protein MGC2718 [Homo sapiens] gb[AAH01076.1]AAH01076 (BC001076) Unknown (protein for MGC:2718) [Homo sapiens] gb|AAH05121.1|AAH05121 (BC005121) 13129046 hypothetical protein MGC2718 [Homo sapiens] 3.2

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
110		gil11466216lrefINP 066539.1l haem lyase [Naegleria	FVALUE
1		gruberi] gb AAG17817.1 AF288092 42 (AF288092)	
2730	11466216	haem lyase [Naegleria gruberi]	1.4
		gi 14701844 gb AAK72251.1  (AF378136) MB2	
2734	14701844	[Plasmodium falciparum]	3.3
2739	7486489	gi 7486489 pir  T00671 hypothetical protein F6E13.4 - Arabidopsis thaliana gb AAC23400.1  (AC004005) putative methyl chloride transferase [Arabidopsis thaliana] gb AAK73255.1  (AY044314) putative methyl chloride transferase [Arabidopsis thaliana]	8.1
2747	7509946	gi 7509946 pir  T26972 hypothetical protein Y47H9C.4- Caenorhabditis elegans emb CAA21739.1  (AL032657) contains similarity to Pfam domain: PF00008 (EGF-like domain), Score=76.2, E-value=2.2e-19, N=17-cDNA EST yk20a5.3 comes from this gene-cDNA EST yk29a12.3 comes from this gene-cDNA EST yk29a12.3 comes from this gene-cDNA EST yk467g8.3 comes > gb AAG60061.1 AF332568_1 (AF332568) CED-1 [Caenorhabditis elegans]	1.6
2749		gi 8569100 gb AAF76445.1 AC015445_12 (AC015445) Contains Ribosomal S17 PF 00366 and DLH PF 01738	0.5
2/49	8309100	domains. [Arabidopsis thaliana] gi 7293625 gb AAF48997.1  (AE003512) CG14223	9.5
2750	7293625	gene product [Drosophila melanogaster]	2.9
2751	5454050	gij5454050 ref NP_006369.1  sema domain, immunoglobulin domain ((g), transmembrane domain ((TM) and short cytoplasmic domain, (semaphorin) 4D; sema domain, immunoglobulin domain (Ig), transmembrane domain (TM) and short cytoplasmic domain, 4D [Homo sapiens] sp(292854 SM4D_HUMAN SEMAPHORIN 4D PRECURSOR (LEUKOCYTE ACTIVATION ANTIGEN CD100) (BB18) (A8) (GR3) gb AAC50810.1  (U60800) semaphorin [Homo sapiens] gi 9964076 gb AAG9812.1 AF275943_1 (AF275943)	9
2761		avermectin polyketide synthase [Streptomyces avermitilis]	2.9
2701	2204070	gi 4234794 gb AAD12962.1  (AF078135) unknown	2.7
2762	4234794	[Leptospira borgpetersenii]	1.3

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	D4-1
SEQ ID	ACCESS	Table 3B Rearest Reignbor (Blast X vs. Non-Redundant	Proteins)
NO NO	N N	DESCRIPTION	P VALUE
		gi 11131078 sp 015072 ATS3 HUMAN ADAM-TS 3	
		PRECURSOR (A DISINTEGRIN AND	
	-	METALLOPROTEINASE WITH	
		THROMBOSPONDIN MOTIFS 3) (ADAMTS-3)	
	]	(ADAM-TS3) dbj BAA20821.1  (AB002364)	
2764	11131078	KIAA0366 [Homo sapiens]	4E-17
	ľ	gi 6449069 gb AAF08806.1 AF192748_1 (AF192748)	
2765	6449069	synapsin Ib [Lampetra fluviatilis]	0.008
	Į.	gi 7300358 gb AAF55517.1  (AE003721) CG8045 gene	
		product [alt 2] [Drosophila melanogaster]	
2769	7300358	gb AAB34837.2  (S78747) RK2 [Drosophila sp.]	3
2772	14731064	gi 14731064 ref XP_036165.1  40679 [Homo sapiens]	6E-20
		gi 4927134 gb AAD33018.1 AF131999 1 (AF131999)	
	l	putative erythrocyte binding protein EBL-1	
2774	4927134		8,6
		gi 11994465 dbi BAB02467.1  (AB025624) contains	
	1	similarity to late embryogenesis abundant	
2775	11994465	protein~gene_id:MLD14.16 [Arabidopsis thaliana]	7.5
		gi 12852706 dbj BAB29508.1  (AK014697) putative	
2776	12852706	[Mus musculus]	1
	ł	-: 0072272 -b A 4 C10/22 1 A G022521 1 (A G022521)	
2777	0072273	gi 9972373 gb AAG10623.1 AC022521_1 (AC022521)  Unknown protein [Arabidopsis thaliana]	3.2
2///	9912313	gill 2838769 dbilBAB24323.1 (AK005931) putative	3.2
2779	12838769	[Mus musculus]	7.6
		gi 14743085 ref XP 050026.1  similar to	
	1	immunoglobulin superfamily containing leucine-rich	
2780	14743085	repeat (H. sapiens) [Homo sapiens]	3.4
	l	gi 4581140 gb AAD24624.1 AC006919_4 (AC006919)	
2781	4581140	unknown protein [Arabidopsis thaliana]	2.8
	1		
		gi 7661684 ref NP_056277.1  DKFZP586L0724 protein	
	1	[Homo sapiens] ref[XP_038194.1  DKFZP586L0724	
		protein [Homo sapiens] ref[XP_038195.1  19734 [Homo	
		sapiens] pir  T14789 hypothetical protein	
		DKFZp586L0724.1 - human cmb CAB53709.1	
	1	(AL110271) hypothetical protein [Homo sapiens]	
2792	7661694	gb AAH01726.1 AAH01726 (BC001726) Similar to	,
2192	/001084	DKFZP586L0724 protein [Homo sapiens] gil13624635[emb]CAA10856.2] (AJ222584) maturase-	7
2793	13624635	[Euglena viridis] [Euglena viridis]	1.3
2193	13027033	mee brown frankring animal	1.3

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESS				
NO	N	DESCRIPTION	P VALUE		
		gi 9622888 gb AAF89968.1 AF200532_1 (AF200532)			
2800	9622888	cellulose synthase-8 [Zea mays]	3.8		
		gi 7499161 pir  T25690 hypothetical protein F15A8.6 -			
		Caenorhabditis elegans gb[AAB52848.1] (U97549)			
		strong similarity to the type-B carboxylesterase/lipase			
2803	7499161	family [Caenorhabditis elegans]	3.1		
		gi 14746401 ref XP 031955.1  ring finger protein 27			
2806	14746401	[Homo sapiens]	0.3		
		gi 14731173 ref XP 017730.2  IQ motif containing			
2809	14731173	GTPase activating protein 2 [Homo sapiens]	2E-11		
		gi 4567214 gb AAD23629.1 AC007113 2 (AC007113)			
2813	4567214	putative villin [Arabidopsis thaliana]	3,5		
		gi 2500866 sp Q20411 SA11 CAEEL SRA-11			
İ		PROTEIN pir T22192 hypothetical protein F44F4.13 -			
		Caenorhabditis elegans emb CAA85461.1  (Z37092)			
ł		contains similarity to Pfam domain: PF02117 (C.elegans			
ļ		Sra family integral membrane protein), Score=675.6, E-			
2817	2500866		3.1		
	200000	gi 15011503 gb AAK77598.1 AF396436_38	3.1		
2818	15011503		5.9		
2010	10011005	(12 556 18 6) yilli i [Testanyillana albimopinia]			
ì		gi 11278033 pir  C81832 transferrin-binding protein A			
l		NMA2024 [imported] - Neisseria meningitidis (group A			
ł		strain Z2491) gblAAC13726.1  (AF058689) transferrin			
ļ		binding protein A precursor [Neisseria meningitidis]			
		emb CAB85243.1  (AL162757) transferrin-binding			
2823	11278033	protein A [Neisseria meningitidis Z2491]	5		
	11270055	protein 11 [110moorie maningman 22491]			
Ì		gi 3913143 sp O23913 AX1B ARATH			
\		ALTERNATIVE OXIDASE 1B PRECURSOR			
		dbj[BAA22624.1] (D89875) alternative oxidase			
1		[Arabidopsis thaliana] dbi BAB01774.1  (AB022215)			
2825	3013142	alternative oxidase 1b precursor [Arabidopsis thaliana]	1.2		
2023	3713143	gi 12842679 dbi BAB25689.1  (AK008476) putative			
2827	12842670	[Mus musculus]	7.8		
2021	12072019	gi 12848636 dbi BAB28031.1  (AK012100) putative	7.0		
2835	12848636	[Mus musculus]	0.083		
2033	12040030	gi 7292152 gb AAF47564.1  (AE003472) Dhc62B gene	0.003		
2839	7292152		16		
2039	1292132	product [Drosopnia meianogaster]	4.6		

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
	ļ.	gi 9628099 ref NP_042686.1  alternative tat protein	
	ļ	[Jembrana disease virus] gb AAA64395.1  (U21603)	
		alternative tat protein [Jembrana disease virus]	
2846	9628099	prf  2116345E tat gene [Jembrana disease virus]	3.3
		gi 4165313 dbj BAA37146.1  (AB022083) SOX30	
2855	4165313	protein [Homo sapiens]	9.2
		gi 14388365 dbj BAB60739.1  (AB062957) hypothetical	
2859	14388365	protein [Macaca fascicularis]	0.55
		gi 14739019 ref XP 005626.3  deleted in bladder cancer	
2864	14720010	chromosome region candidate 1 [Homo sapiens]	6.3
2804	14/39019		6.3
	l	gi 6319950 ref NP_010031.1  Transcription regulator;	
		Ycr106wp [Saccharomyces cerevisiae]	
		sp P25611 YCZ6_YEAST PUTATIVE 95.7 KD	
		TRANSCRIPTIONAL REGULATORY PROTEIN IN	
	ļ	PAU3-AAD3 INTERGENIC REGION pir   S19418	
		probable membrane protein YCR106w - yeast	
		(Saccharomyces cerevisiae) emb CAA42238.1	
20.67	C210050	(X59720) hypothetical protein [Saccharomyces	0.1
2867	6319950	cerevisiae]	9.1
2870	14724850	gi 14724850 ref XP 050192.1  29140 [Homo sapiens]	2.2
2070	11721000	gi 4587097 dbi BAA76616.1  (AB019045)	
2871	4587097	OMPdecarboxylase [Rhizomucor pusillus]	9.7
		gi 7522108 pir  T29097 pro-pol-dUTPase polyprotein -	
	1	murine endogenous retrovirus ERV-L (fragment)	
	i	emb CAA73251.1  (Y12713) protease; reverse	
		transcriptase; RNaseH; integrase; dUTPase; Pro-Pol-	
2873	7522108	dUTPase polyprotein [Mus musculus]	0.003
		gi 7482073 pir  B69010 conserved hypothetical protein	
	Į.	MTH1078 - Methanobacterium thermoautotrophicum	
	1	(strain Delta H) gb AAB85567.1  (AE000879)	
		conserved protein [Methanothermobacter	
2875	7482073	thermautotrophicus]	4.6
2876	11360401		0.000000005
		gi 5091521 dbj BAA78756.1  (AB023482) Hypothetical	-
2879	5091521	protein [Oryza sativa]	2.1_
		gi 13810543 dbj BAB43950.1  (AB051633) ookinete	
2881	13810543	surface protein Pos28-2 [Plasmodium ovale]	3
		gi 5091521 dbj BAA78756.1  (AB023482) Hypothetical	
2882	5091521	protein [Oryza sativa]	2.1

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N -	DESCRIPTION	P VALUE
		gi 6723244 dbj BAA89640.1  (AB036666) similar to	
1		terminase large subunit of phage lambda [Wolbachia sp.	
2887	6723244	wKue]	7.1
		gi 7494347 pir  E71625 variant-specific surface protein	
ì		1 truncated homolog PFB0020c - malaria parasite	
		(Plasmodium falciparum) gb AAC71794.1  (AE001366)	
2889	7494347	PfEMP1 fragment [Plasmodium falciparum]	5
		gi 6942188 gb AAF32349.1 AF220008_1 (AF220008)	
2890	6942188	coilin p80 [Danio rerio]	0.11
		gi 10639397 emb CAC11399.1  (AL445063)	
	l	hypothetical membrane protein [Thermoplasma	
2891	10639397	acidophilum]	8.5
		gi 10800417 ref NP_006759.2  BRCA1 associated	
2898	10800417	protein [Homo sapiens]	0.98
		gi 7488089 pir  T02283 probable disease resistance	
		protein T13D8.20 - Arabidopsis thaliana	
		gb AAC24071.1  (AC004473) Contains similarity to	
		TMV resistance protein N homolog gb 2245048 from A.	
2901	7488089	thaliana chromosome 4 contig gb Z97342. [Arabidopsis thaliana]	8.5
2901	7400009	inananaj	0.3
		gi 10728394 gb AAF45654,2  (AE003421) EG:171E4.2	
2904	10728394	gene product [Drosophila melanogaster]	1.4
2501	10720331	gil10956333 refINP 052782.1  pXO1-86 [Bacillus	2.1
		anthracis] pir  F59101 hypothetical protein pXO1-86 -	
		Bacillus anthracis virulence plasmid pXO1	
	l	gb AAD32390.1 AAD32390 (AF065404) pXO1-86	·
2905	10956333	[Bacillus anthracis]	9.7
		gi 4028153 gb AAC96117.1  (AF083221) putative	
2906	4028153	ncurotransmitter receptor [Takifugu rubripes]	2.8
		gi 12836120 dbj BAB23511.1  (AK004729) putative	
2909	12836120	[Mus musculus]	5.3
		gi 1778844 gb AAB40929.1  (U83086) LimA	
2911	1778844	[Dictyostelium discoideum]	1.3
		gi 3192956 gb AAC41298.1  (AF033670) T-Box protein	
2913	3192956	4 [Gallus gallus]	3.1
		gi 7506147 pir  T33303 hypothetical protein R01B10.4 -	
2014	7506145	Caenorhabditis elegans gb AAC17768.1  (AF068718)	
2914	7506147	R01B10.4 gene product [Caenorhabditis elegans]	0.4

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#### Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins) SEQ ID ACCESS NO N DESCRIPTION P VALUE gi|6041793|gb|AAF02113.1|AC009755 6 (AC009755) putative auxin-independent growth promoter 2915 6041793 [Arabidopsis thaliana] 2.1 gil5803098|refINP 006757.1| zinc finger protein 220: Monocytic leukemia zinc finger protein [Homo sapiens] sp|Q92794|MOZ HUMAN MONOCYTIC LEUKEMIA ZINC FINGER PROTEIN (ZINC FINGER PROTEIN 220) gb(AAC50662.11 (U47742) 2916 5803098 monocytic leukaemia zinc finger protein [Homo sapiens] 0.084 gil10639353lemblCAC11355.11 (AL445063) hypothetical membrane protein [Thermoplasma 2919 10639353 acidophilum] 3.1 gil 13235586|emb|CAC33776.1| (AJ301807) SclB 2920 13235586 protein [Streptococcus pyogenes] 9.7 gi|7507618|pir||T33548 hypothetical protein T10D4.8 -2925 7507618 Caenorhabditis elegans 4.3 gi|7522108|pirl|T29097 pro-pol-dUTPase polyprotein murine endogenous retrovirus ERV-L (fragment) emb|CAA73251.1| (Y12713) protease; reverse transcriptase; RNaseH; integrase; dUTPase; Pro-Pol-2928 7522108 dUTPase polyprotein [Mus musculus] 1.2 gi|13542796|gb|AAH05601.1|AAH05601 (BC005601) Similar to RIKEN cDNA 1110061A19 gene [Mus 2932 13542796 musculus) 6.4 gil417869|sp|P33007|TERP PSESP TERPREDOXIN (TDX) pir E42971 terpredoxin - Pseudomonas sp gb|AAA25998.1| (M91440) terpredoxin [Pseudomonas 2934 417869 3.7 gi|7433891|pir||T00981 flavonol 3-O-glucosyltransferase homolog T9J22.15 - Arabidopsis thaliana gb|AAC14497.1| (AC002505) putative 2938 7433891 glucosyltransferase [Arabidopsis thaliana] 6.8 gi|15021546|gb|AAK77823.1|AF369029 154 2939 15021546 (AF369029) ORF154 [white spot syndrome virus] 0.25 gi|11466694|refINP 039290.1| ORF370i [Marchantia polymorphal splP12174lMATK, MARPO PROBABLE INTRON MATURASE pir A05034 hypothetical protein 370i - liverwort (Marchantia polymorpha) chloroplast cmb|CAA28076.1| (X04465) ORF370i 11466694 [Marchantia polymorpha] 4.3

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 7302236 gb AAF57330.1  (AE003786) CG10416	
2945	7302236	gene product [Drosophila melanogaster]	1.4
		gi 2495638 sp P76511 YFDO_ECOLI	
	1	HYPOTHETICAL 14.4 KDA PROTEIN IN INTC-	
		DSDC INTERGENIC REGION pir  C65009	
		hypothetical protein b2358 - Escherichia coli (strain K-	
		12) gb AAC75417.1  (AE000324) orf, hypothetical	
2950	2495638	protein [Escherichia coli K12]	4.5
		gi 220578 dbj BAA00447.1  (D00570) open reading	
2955	220578	frame (251 AA) [Mus musculus]	4.2
	1	gi 12855573 dbj BAB30384.1  (AK016695) putative	
2958	12855573		0.007
		gi 2981631 dbj BAA25253.1  (AB012223) ORF2 [Canis	
2963	2981631	familiaris]	0.29
	1		
		gi 7657401 ref NP_056616.1  neuropathy target	
		esterase; Swiss cheese [Mus musculus]	
		gb AAD51700.1 AF173829_1 (AF173829) neuropathy	
2966	7657401	target esterase homolog [Mus musculus]	1.5
		gi 14285535 sp P71399 LSG1_HAEIN LSG LOCUS	
2967	14285535	PUTATIVE PROTEIN 1	4.8
		gi 14423780 sp 095013 04F3_HUMAN OLFACTORY	
		RECEPTOR 4F3 gb AAD05195.1  (AC004908) similar	
		to rat olfactory receptor OR18; similar to S29710	
2973	14423780	(PID:g423702) [Homo sapiens]	3E-35
2975	14771601	gi 14771691 ref XP 045484.1  67354 [Homo sapiens]	1.2
2913	14//1091	gi 14771091 1et XF_045464.1  07354 [Hollo saptens]	1.3
2978	6573738	F20B24.13 [Arabidopsis thaliana]	8.5
2976	03/3/38	gi 13811437 gb AAK40121.1  (AF354707) type II	0.3
2982	13811437	deodinase [Xenopus laevis]	2.4
2702	13011437	deodinase [Actiopus taevis]	2.4
1		-: 7522109 ::: F7200071 HFTP	
		gi 7522108 pir  T29097 pro-pol-dUTPase polyprotein - murine endogenous retrovirus ERV-L (fragment)	
		emb CAA73251.1  (Y12713) protease; reverse	
		transcriptase; RNascH; integrase; dUTPase; Pro-Pol-	
2986	7522108	dUTPase polyprotein [Mus musculus]	0.00001
2200	7322106	gi 12722927 gb AAK04181.1 AE006247 3 (AE006247)	0.00001
		UNKNOWN PROTEIN [Lactococcus lactis subsp.	
2990	12722927	lactisl	1.3
2330	LILLIZI	monaj	1.3

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WC0214500 [flig //E /WO0214500 opc]

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	The section of the se	1 Totellis)
NO	N	DESCRIPTION	P VALUE
		gi 11346713 pir  F81302 hypothetical protein Ci1009c	
		[imported] - Campylobacter jejuni (strain NCTC 11168)	
		emb CAB73265.1  (AL139077) hypothetical protein	
2994	11346713	Cj1009c [Campylobacter jejuni]	4.1
		URDO 10 451 114 4 GCOSO 4 BLOCK CORES	
2995	7321945	gi[7321945 gb AAC60504.2  (S68356) action potential broadening potassium channel [Aplysia sp.]	4.0
2993	1321943	gi 11360394pir  T42731 atrophin-1 related protein - rat	4.2
		gb AAA98970.1  (U44091) atrophin-1 related protein - rat	
3003	11360304	[Rattus norvegicus]	7.6
3003	11300394	gi 10765337 gb AAG22997.1  (AF188579) glycoprotein	7.0
3005	10765337	[Bovine respiratory syncytial virus]	1.9
3003	10103331	gi 5803252 dbi BAA83562.1  (AP000399) hypothetical	1.9
3006	5803252	protein [Oryza sativa]	0.046
	0000202	gi 7662688 gb AAF66138.1  (L00016) urf4 [Homo	0.040
3007	7662688	sapiens	1.7
		gi 14150037 ref NP 115666.1  hypothetical protein	
		DKFZp761C121 [Homo sapiens] refiXP 027894.1	
		hypothetical protein DKFZp761C121 [Homo sapiens]	
		emb CAB66495.1  (AL136560) hypothetical protein	
3013	14150037	[Homo sapiens]	0.0000001
		- 170 4070 ( - 1-1-1 A P 40 40 1 11 A P 20 20 0 1 1 ( A P 20 20 0 1 )	
3014	7242706	gi 7243706 gb AAF43421.1 AF233291_1 (AF233291) epsin-like protein [Drosophila melanogaster]	6.5
3014	1243100	gi 7293274 gb AAF48655.1  (AE003503) CG9644 gene	6.3
3019	7202274	product [Drosophila melanogaster]	0.47
3019	1233214	gi 11414881 dbj BAB18568.1  (AB028173) HCCA2	0.47
3025	11414881	[Homo sapiens]	8.6
		gi 11359423 pir  T48729 hypothetical protein 8D4.30	
		[imported] - Neurospora crassa emb CAB88545.1	
		(AL353819) conserved hypothetical protein	
3026	11359423	[Neurospora crassa]	0.66
		gi 2118405 pir  I51018 cobra venom factor precursor -	
		monocled cobra gb[AAA68989,1] (U09969) cobra	
3028	2118405	venom factor precursor [Naja naja]	3.1
	2110103	gi 2739145 gb AAC98522.1  (AF030306) envelope	5.1
		protein; ORF4 [Porcine reproductive and respiratory	
3029	2739145		4.3
		gi 13549158 gb AAK29672.1 AF353095 1 (AF353095)	
		protein synthesis initiation factor eIF2 beta [Arabidopsis	
3030	13549158		6.3

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
3034	14753542	gi 14753542 ref XP_046011.1  53288 [Homo sapiens]	0.000000006
3042	3150471	gi 3150471 gb AAC16989.1  (AF067211) Hypothetical protein B0205.2 [Caenorhabditis elegans]	0.56
3044	14734012	gi 14734012 ref XP_051005.1  KIAA1297 protein [Homo sapiens]	0.16
3047	7493994	gi 7493994 pir  IC6564 cellobiose oxidase (EC 1.1.3.25) precursor - white-rot fungus (Trametes versicolor)	6.1
3056	114351	gi 114351 sp P08314 ATI2_HSVIF ALPHA TRANS- INDUCING FACTOR 77 KD PROTEIN pir  TNBE77 77K alpha trans-inducing protein - human herpesvirus 1 (strain F) gb AAA45768.1  (M15621) alpha trans- inducing factor 77kb [human herpesvirus 1]	7
3069		gi 6562755 emb CAB62894.1  (AL035475) possible ribosomal protein [Plasmodium falciparum]	6.4
3071	804764	gi 804764 gb AAA65999.1  (M27826) neutral protease large subunit [Homo sapiens]	0.00003
3079	14768311	gi 14768311 ref[XP_048396.1  methyl CpG binding protein 2 [Homo sapiens]	0.65
3080	11994604	gi 11994604 dbj BAB02658.1  (AP002062) gene_id:T22B15.11~unknown protein [Arabidopsis thaliana] gi 7302673 gb AAF57753.1  (AE003800) stau gene	5
3083	7302673	product [alt 2] [Drosophila melanogaster]	9.2
3085	7494170	gij/1494170 pir  D71613 GAF domain protein (cyclic nt signal transduct, PFB0510w - malaria parasite (Plasmodium falciparum) gb AAC71891.1  (AE001399) GAF domain protein (cyclic nt signal transduct.) [Plasmodium falciparum]	8.1
3087	14590168	gi 14590168 ref NP_142233.1  hypothetical protein [Pyrecoccus horikoshii] pir [F/1247 hypothetical protein PH0237 - Pyrococcus horikoshii dbj BAA29309.1  (AP000001) 230aa long hypothetical protein [Pyrococcus horikoshii]	2.3
3097	14326099	gi 14326099 gb AAK60138.1 AF365405_1 (AF365405) ribosomal protein S14 [Schizosaccharomyces pombe]	3

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundam	Di
SEQ ID	ACCESS	Table 36 Nearest Neighbor (BlastX Vs. Non-Redundam	Proteins)
NO NO	N N	DESCRIPTION	P VALUE
		gi 2499244 sp Q35140 NU2M NEUCR NADH-	
	1	UBIQUINONE OXIDOREDUCTASE CHAIN 2	
		pir  A25096 NADH dehydrogenase (ubiquinone) (EC	
	1	1.6.5.3) chain 2 - Neurospora crassa mitochondrion	
	(	emb CAA27418.1  (X03793) put. URF-2 like protein	
3098	2499244	(aa 1-583) [Neurospora crassa]	8.1
	1	gi 113668 sp P23961 ALUC_HUMAN !!!! ALU	
3101	113668	CLASS C WARNING ENTRY !!!!	6.5
		gi 12721131 gb AAK02908.1  (AE006121) unknown	
3105	12721131	[Pasteurella multocida]	9.2
	1	gi 12621134 ref NP_075244.1  MEGF6 [Rattus	
	1	norvegicus] pir  T13954 MEGF6 protein - rat	}
0107		dbj BAA32462.1  (AB011532) MEGF6 [Rattus	
3107	12621134	norvegicus]	2.1
		gi 1346666 sp P48305 NB5M_BOVIN NADH-	
	-	UBIQUINONE OXIDOREDUCTASE B15 SUBUNIT	
		(COMPLEX I-B15) (CI-B15) pir  S28237 NADH	
	1	dehydrogenase (ubiquinone) (EC 1.6.5.3) chain B15 -	
		bovine emb CAA46107.1  (X64898) B15 subunit of the NADH: ubiquinone oxidoreductase complex [Bos	
3108	1346666		0.47
3100	1340000	<del></del>	0.47
		gi 462022 sp P33948 ERD2_PLAFA ER LUMEN	
	1	PROTEIN RETAINING RECEPTOR pir  S39609 ERD2 protein - malaria parasite (Plasmodium	
	i	falciparum) emb CAA81128.1  (Z26043) ERD2	
	ŀ	[Plasmodium falciparum] emb[CAA52861,1] (X74869)	
3111	462022	PFERD2 [Plasmodium falciparum]	0.95
		gi 12831427 gb AAK02082.1  (AY014401) site-specific	
3114	12831427	recombinase IntIA [Listonella pelagia]	8.3
	1	gi 7657530 ref NP_055248.1  rhabdoid tumor deletion	
		region protein 1 [Homo sapiens] ref[XP_009866.1]	
	Į	rhabdoid tumor deletion region protein 1 [Homo	
	1	sapiens] ref[XP_037053.1  rhabdoid tumor deletion	
		region protein 1 [Homo sapiens]	
	ĺ	gb AAF02484.1 AF133587_1 (AF133587) rhabdoid	
		tumor deletion region protein 1 [Homo sapiens]	
2117	7657520	gb AAH08986.1 AAH08986 (BC008986) rhabdoid	7.0
3117	7657530	tumor deletion region protein 1 [Homo sapiens]	7.8
		gi 12083527 gb AAG48836.1 AC084218_6	
		(AC084218) similar to Arabidopsis thaliana DNA-	
3123	12092527	directed RNA polymerase (EC 2.7.7.6) II largest chain (JDMU1) [Oryza sativa]	6.9
3123	12003327	(IDMO1) [Oryza sauvaj	0.9

	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS			
NO	N	DESCRIPTION	P VALUE	
		gi 2565196 gb AAB81938.1  (AF000381) non-functional		
3126	2565196	folate binding protein [Homo sapiens]	8.8	
		gi 8923808 ref NP_060941.1  uncharacterized		
	{	hypothalamus protein HT010 [Homo sapiens]		
	i	gb AAF67649.1 AF220184_1 (AF220184)		
	}	uncharacterized hypothalamus protein HT010 [Homo		
3132	8923808	sapiens]	7E-13	
	1	gi 14755072 ref XP_048449.1  hypothetical protein		
3138	14755072	XP_048449 [Homo sapiens]	9.5	
ļ		gi 1085507 pir  S52306 zinc finger protein 10 - mouse		
		emb CAA85283.1  (Z36270) GC Binding Protein - 23b		
3141	1085507	[Mus musculus]	6.7	
Į į	l	gi 4417278 gb AAD20403.1  (AC007019) unknown		
3142	4417278	protein [Arabidopsis thaliana]	2.8	
		gi 12084884 ref[NP_073349.1  UL54 post-translational		
		gene regulation protein [Meleagrid herpesvirus 1]		
1	ŀ	gb AAG45793.1 AF291866_60 (AF291866) UL54 post-		
		translational gene regulation protein [Meleagrid		
3148	12084884	herpesvirus 1]	8.2	
	1	gi 13122204 emb CAB89584.2  (AL354512) possible		
		non-canonical ubiquitin conjugating enzyme 1		
3151	13122204	[Leishmania major]	0.65	
	1			
1	1	gi 418745 pir  S34959 NADH dehydrogenase		
	1	(ubiquinone) (EC 1.6.5.3) chain 4 - Crithidia oncopelti		
		mitochondrion emb CAA39491.1  (X56015) NADH		
3153	418745	dehydrogenase subunit 4 [Crithidia oncopelti]	1	
		gi 6015766 emb CAB57593.1  (Y18930) hypothetical		
3157	6015766	protein [Sulfolobus solfataricus]	7.7	
	1	gi 6983867 dbj BAA90802.1  (AP001168) Similar to		
2162	6002067	putative salt-inducible protein (AC006248) [Oryza	2.0	
3162	6983867	sativa]	2.8	
	1			
1		gi 11467948 sp 065404 ER11_ARATH SQUALENE		
1	1	MONOOXYGENASE 1,1 (SQUALENE EPOXIDASE		
3163	11467040	1,1) (SE 1,1) dbj BAB08406.1  (AB016883) squalene monooxygenase [Arabidopsis thaliana]	6.1	
3103	1140/948		0.1	
	1	gi]3461813 gb AAC32907.1  (AC004138) putative		
	1	sucrose-proton symporter [Arabidopsis thaliana] emb[CAB92307.1] (AJ289165) sucrose transporter		
3167	3461813	[Arabidopsis thaliana]	4.4	
3107	3401013	[/Maninobara manana]	4.4	

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
-10		DESCRIPTION	FVALUE
		gi 6753576 ref NP 034128.1  cytochrome P450, 2b10,	
	1	phenobarbitol inducible, type b [Mus musculus]	
	i	sp P12791 CPBA MOUSE CYTOCHROME P450	ĺ
		2B10 (CYPIIB10) (TESTOSTERONE 16-ALPHA	
	1	HYDROXYLASE) (P450-16-ALPHA) (CLONE	
1		PF3/46) pir  B31047 testosterone 16alpha-hydroxylase	1
		(EC 1.14.14) cytochrome P450 2B10 - mouse	
İ		gb AAA40425.1  (M21856) testosterone 16-alpha-	
3168	6753576	hydroxylase [Mus musculus]	7.6
		Second - Occasioner Course Della Strong	
		gi 585811 sp Q08517 REFR_SPVKA RIFAMPICIN  RESISTANCE PROTEIN (62 KD PROTEIN)	}
3174	585811	gb AAA16176.1  (L22012) ORF H1L [Swinepox virus]	6.9
3174	303011	gojaaarot 70.1 (E22012) OAF ATE [Switepox virus]	0.9
		gi 14714688 gb AAH10485.1 AAH10485 (BC010485)	
3176	14714688	Unknown (protein for MGC:7224) [Mus musculus]	2.1
		gi 10047191 dbj BAB13389.1  (AB046783) KIAA1563	
3178	10047191	protein [Homo sapiens]	4E-26
		gi 6324246 ref NP_014316.1  Ynl083wp	
		[Saccharomyces cerevisiae] pir  S57539 probable	
	1	membrane protein YNL083w - yeast (Saccharomyces	
1		cerevisiae) emb CAA61427.1  (X89016) ORF N2312	
2100	620.40.46	[Saccharomyces cerevisiae] emb CAA95958.1	0.24
3180	6324246	(Z71359) ORF YNL083w [Saccharomyces cerevisiae]	0.24
	1	gi 7494290 pir  C71618 hypothetical protein PFB0315w malaria parasite (Plasmodium falciparum)	
1		gb AAC71852.1  (AE001386) hypothetical protein	
3184	7494290	[Plasmodium falciparum]	0.084
	7.15.155.0	[Financount Autoput un]	0.001
		gi 6754948 ref NP 036089,1  origin recognition	
		complex, subunit 5 homolog (S. cerevisiae); mouse	
		origin recognition complex 5 [Mus musculus]	
		sp Q9WUV0 ORC5 MOUSE ORIGIN	
		RECOGNITION COMPLEX SUBUNIT 5	
		emb CAB43767.1  (AJ007360) ORC5-related protein	
		[Mus musculus] gb AAH06927.1 AAH06927	
		(BC006927) origin recognition complex, subunit 5	
3185		homolog (S. cerevisiae) [Mus musculus]	8.5
2100		gi 7295489 gb AAF50803.1  (AE003567) CG10671	4.2
3199	7295489	gene product [Drosophila melanogaster]	4.3

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DEGGDTDWYON	
NU	I N	DESCRIPTION gi 14741894 ref XP_049656.1  hypothetical protein	P VALUE
3203	14741894	XP_049656 [Homo sapiens]	4
		gi 14520989 ref NP 126464.1  hypothetical protein	ļ
1		[Pyrococcus abyssi] pir  F75122 hypothetical protein	
1	l	PAB1844 - Pyrococcus abyssi (strain Orsay)	
ł	1	emb CAB49695.1  (AJ248285) hypothetical protein	
3208	14520989	[Pyrococcus abyssi]	4.1
1		gi 9911016 sp P76543 YFFL_ECOLI	
	1	HYPOTHETICAL 25.1 KDA PROTEIN IN EUTB-	
}		EUTH INTERGENIC REGION pir    B65019	
		hypothetical protein b2443 - Escherichia coli (strain K- 12) gblAAC75496.1 (AE000331) orf, hypothetical	
3210	9911016	protein [Escherichia coli K12]	7.9
- <del></del>		gi 9626990 ref NP 056868.1  No definition line found	
ł		gb AAB92648.1  (AF035403) No definition line found	
3211	9626990		3.7
1			
1		gi 7504416 pir  T16477 hypothetical protein F56D2.5 -	
2212	2504416	Caenorhabditis elegans gb AAB52683.1  (U13644)	
3213	7504416	F56D2.5 gene product [Caenorhabditis elegans] gi[14746120 ref XP 005163.2  cadherin 17, LI cadherin	8,4
l		(liver-intestine) [Homo sapiens] ref[XP 043524.1]	
ł		cadherin 17, LI cadherin (liver-intestine) [Homo	
3216	14746120		1E-18
l		gi 9366835 emb CAB95597.1  (AL359782) hypothetical	
3217	9366835	protein, CHR1.368 [Trypanosoma brucei]	4.1
1		Transport III A Porogo VIA Porogo II -	
3218	6272295	gi 6272295 gb AAF06072.1 AF072715_3 (AF072715) unknown [Mycoplasma mycoides subsp. capri]	0.17
3216	0212293	gi 7293488 gb AAF48863.1  (AE003509) CG15040	0.17
3220	7293488	gene product [Drosophila melanogaster]	2.8
		gi 7662432 ref NP_055769.1  KIAA0985 protein [Homo	
1		sapiens] ref[XP_007006.3  KIAA0985 protein [Homo	
		sapiens] sp Q9Y2J0 RP3A_HUMAN RABPHILIN-3A	
		dbj BAA76829.1  (AB023202) KIAA0985 protein	
3223	7662432	[Homo sapiens]	0.18
1		gi 7504860 pir  T33299 hypothetical protein H05B21.4 - Caenorhabditis elegans gb AAC17764.1  (AF068717)	
ļ		Hypothetical protein H05B21.4 [Caenorhabditis	
3225	7504860	elegans]	3.7
		······································	

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L		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESS					
NO	N	DESCRIPTION	P VALUE			
		gi 6708056 gb AAF25780.1 AF163311_1 (AF163311)				
3226	6708056	Kall.2 [Danio rerio]	2.5			
l	ſ	gi 13473697 ref NP_105265.1  acyl-CoA thioesterase				
		[Mesorhizobium loti] dbj BAB51051.1  (AP003004)				
3229	13473697	acyl-CoA thioesterase [Mesorhizobium loti]	5.4			
1	1	gi 12964703 gb AAK11280.1 AF315035_1 (AF315035)				
		phosphocholine cytidylyltransferase [Plasmodium				
3230	12964703	falciparum]	2.1			
		gi 7521951 pir  T30180 hypothetical protein -				
l	l	Shewanella sp gb AAB81120.1  (U73935) unknown				
3231	7521951		6			
		gi 3249620 gb AAC24120.1  (AF067182) IDI-2				
3232	3249620	precursor [Podospora anscrina]	3.1			
		gi 7493160 pir  T40507 probable vacuolar protein				
		sorting-associated pro tein - fission yeast				
1	İ	(Schizosaccharomyces pombe) emb[CAA20730.1]				
		(AL031534) putative vacuolar protein sorting-				
3233	7493160	associated protein [Schizosaccharomyces pombe]	5.5			
		gi 7462822 pir  C72269 hypothetical protein TM1297 -				
		Thermotoga maritima (strain MSB8)				
		gb AAD36371.1 AE001785_2 (AE001785)				
3237	7462822	oxidoreductase, putative [Thermotoga maritima]	5.5			
		gi 1531651 gb AAC61662.1  (U67083) KRAB-zinc				
3241	1531651	finger protein KZF-2 [Rattus norvegicus]	4.6			
1	1	gi 2118405 pir  I51018 cobra venom factor precursor -				
		monocled cobra gb AAA68989.1  (U09969) cobra				
3247	2118405	venom factor precursor [Naja naja]	9.2			
1	1	gi 11499508 ref NP_070749.1  LSU ribosomal protein				
		L4P (rpl4P) [Archaeoglobus fulgidus]				
		sp O28355 RL4_ARCFU 50S RIBOSOMAL				
1	1	PROTEIN L4/L1E pir C69490 LSU ribosomal protein				
		L4P (rpl4P) homolog - Archaeoglobus fulgidus				
		gb AAB89332.1  (AE000971) LSU ribosomal protein				
3253	11499508	L4P (rpl4P) [Archaeoglobus fulgidus]	2.9			

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
110		DESCRIPTION	1 VALUE
ĺ			
1	1	gi 461509 sp Q03045 AMYG HORRE	
1		GLUCOAMYLASE P PRECURSOR (GLUCAN 1.4-	
ļ.		ALPHA-GLUCOSIDASE) (1,4-ALPHA-D-GLUCAN	
		GLUCOHYDROLASE) pir S33908 glucan 1,4-alpha-	
l		glucosidase (EC 3.2.1.3) P precursor - creosote fungus	
1		cmb CAA47945.1  (X67708) 1, 4-alpha-D-glucan	
1	ļ	glucohydrolase; glucan 1,4-alpha-glucosidase	
ì	ĺ	[Amorphotheca resinae] emb CAA48243.1  (X68143)	
ŀ		glucan 1,4-alpha-glucosidase [Amorphotheca resinae]	
		prf  1907167A glucoamylase P [Amorphotheca resinae]	
3258	461509	prf  2113213A glucoamylase P [Amorphotheca resinae]	5,8
		-359170001-1-14 AD52074 11 (AD174771)	
3260	5017000	gi 5817900 gb AAD52976.1  (AF176771) reverse transcriptase [Human immunodeficiency virus type 2]	٠,
3260	3817900	transcriptase [Human immunodenciency virus type 2]	6.1
		gi 12804147 gb AAH02928.1 AAH02928 (BC002928)	
3262	12804147	Unknown (protein for MGC:11357) [Homo sapiens]	3E-14
3202	12004147	Chikhowh (protein for MGC.11337) [Homo sapiens]	3E-14
		-: 6222756 (DVID_012827_1	
1	ì	gi 6323756 ref NP_013827.1  myosin I; Myo5p  Saccharomyces cerevisiael sp 004439 YMZ9 YEAST	
		HYPOTHETICAL MYOSIN-LIKE PROTEIN IN	
1		ILV2-ADE17 INTERGENIC REGION pir S54570	
		probable membrane protein YMR109w - yeast	
		(Saccharomyces cerevisiae) emb CAA89745.1	
		(Z49702) unknown [Saccharomyces cerevisiae]	
		gb[AAB37419.1] Myo5p=actin patch localized myosin I	
1		variant/MYO5 product [Saccharomyces	
3266	6323756	cerevisiae=yeast, Peptide, 1219 aa]	0.24
		gi 12848866 dbj BAB28115.1  (AK012246) putative	
3271	12848866	[Mus musculus]	0.011
		gi 4758842 ref NP_004551.1  receptor tyrosine kinase-	
1	ĺ	like orphan receptor 2; Ror2; neurotrophic tyrosine	
1		kinase, receptor-related 2 [Homo sapiens]	
Į.	l	sp Q01974 ROR2_HUMAN TYROSINE-PROTEIN	
		KINASE TRANSMEMBRANE RECEPTOR ROR2	
		PRECURSOR (NEUROTROPHIC TYROSINE	
1		KINASE, RECEPTOR-RELATED 2) pir B45082	
ì	)	neurotrophic receptor ror2 precursor - human	
		gb AAA60276.1  (M97639) transmembrane receptor	
3273	4758842	[Homo sapiens]	5.6

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3293

3295

3301

6

0.16

8.3

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#### Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins) SEQ ID ACCESS NO N DESCRIPTION P VALUE gi|913140|gb|AAB33486.1| ARK2 product/receptor-like serine/threonine protein kinase ARK2 [Arabidopsis 3274 913140 thaliana, Columbia, Pentide, 850 aa1 1.7 gi|11357726|pir||T51437 hypothetical protein F2G14 40 - Arabidopsis thaliana emb|CAC01811.1| (AL391146) putative protein [Arabidopsis thaliana] gb|AAK49610.1|AF372894 1 (AF372894) AT5g14920/F2G14 40 [Arabidopsis thaliana] gb|AAK74054.1| (AY045696) AT5g14920/F2G14 40 3276 11357726 [Arabidopsis thaliana] 2 gil7511204|pir||T27899 hypothetical protein ZK546.5 -Caenorhabditis elegans gb|AAA68738.1| (U29380) 3277 7511204 Hypothetical protein ZK546.5 [Caenorhabditis elegans] 3.4 gi|7492154|pir||T38426 major facilitator protein 3279 7492154 homolog - fission yeast (Schizosaccharomyces pombe) 6 gil14764251|refIXP 011989.3| ELKL motif kinase 1 3285 14764251 isoform a [Homo sapiens] 9E-18 gi|126686|sp|P27424|MGP1 BOVIN MICROFIBRIL-ASSOCIATED GLYCOPROTEIN PRECURSOR (MAGP) (MAGP-1) (TROPOELASTIN-BINDING PROTEIN) pir||A54151 microfibril-associated glycoprotein precursor - bovine gb[AAA62715.1] (M59851) microfibril-associated glycoprotein [Bos taurus] gb|AAB29686.1| (S68064) microfibrilassociated glycoprotein, MAGP=tropoelastin-binding 3286 126686 protein [cattle, Peptide Partial, 183 aal [Bos taurus] 5.4 gi|13622462|gb|AAK34181.1| (AE006573) conserved

13622462 hypothetical protein [Streptococcus pyogenes M1 GAS]

5815436 98kDa HDM allergen [Dermatophagoides farinae]

7523494 protein [Oryza sativa]

gi|7523494|dbi|BAA94222.1| (AP001633) hypothetical

gi|5815436|gb|AAD52672.1|AF178772 1 (AF178772)

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ano m		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
		gi 7522146 pir  T17467 rifamycin polyketide synthase	
		modules 9-10 - Amycolatopsis mediterranei	
		emb CAA11039.1  (AJ223012) rifamycin polyketide	
		synthase, type 1 [Amycolatopsis mediterranci]	
		gb AAC01714.1  (AF040570) polyketide synthase	
3302	7522146	[Amycolatopsis mediterranei]	8.5
		gi 14737876 ref XP_031833.1  collagen, type V, alpha 1	
3313	14737876	[Homo sapiens]	9.6
		Hanna con the control of the control	
ĺ		gi 10092673 ref NP_064709.1  hypothetical protein	
	l	[Homo sapiens] ref[XP_048065.1  hypothetical protein [Homo sapiens] gb[AAC24312.1  (AC004382)	
3316	10002672	Unknown gene product [Homo sapiens]	0.000002
3310	10092073		0.000002
		gi 135937 sp P04924 TNFA_RABIT TUMOR NECROSIS FACTOR PRECURSOR (TNF-ALPHA)	
	Ì	(CACHECTIN) gb AAA31482.1  (M12846) tumor	
		necrosis factor [Oryctolagus cuniculus]	
}	<b>\</b>	gb[AAA31484.1] (M60340) tumor necrosis factor	
3317	135937	[Oryctolagus cuniculus]	8.2
	100501	[Or) Transpar Tameuras	0.2
		gi 11282039 pir  C82096 aminoacyl-histidine dipeptidase	
		VC2279 [imported] - Vibrio cholerae (group O1 strain	
		N16961) gb AAF95423.1  (AE004299) aminoacyl-	
3318	11282039	histidine dipeptidase [Vibrio cholerae]	0.73
1	i '	gi 11360605 pir  A81752 conserved hypothetical protein	
ł I	Į į	TC0009 [imported] - Chlamydia muridarum (strain	
		Nigg) gb AAF38902.1  (AE002269) conserved	
3320	11360605	hypothetical protein [Chlamydia muridarum]	9.2
		gi 7491033 pir  T38495 hypothetical protein	
		SPAC29B12.07 - fission yeast (Schizosaccharomyces	
3321	7491033	pombe)	0.46
		gi 102425 pir  B41132 collagen-related protein 2 - Hydra	
		magnipapillata (fragment) pir  S21930 mini-collagen -	
2222		Hydra sp emb CAA43380.1  (X61046) mini-collagen	0.00
3322	102425	[Hydra sp.]	0.99
		-i/7507644 -i-i/T25997	
		gi 7507644 pir  T25887 hypothetical protein T10H10.2 - Caenorhabditis elegans gb AAB37989.1  (U80848)	
3328	7507644	T10H10.2 gene product [Caenorhabditis elegans]	4.3
5520	7507044	1101110.2 gene product [Cacitornabulus cicgans]	7.3

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS			
NO	N	DESCRIPTION	P VALUE	
		gi 3738325 gb AAC63666.1  (AC005170) cyclic		
		nucleotide and calmodulin-regulated ion channel		
3331	3738325	[Arabidopsis thaliana]	9.1	
1	<b>\</b>			
	i	gi 8978523 dbj BAA98360.1  (AP002545) CT147		
3332	8978523	hypothetical protein [Chlamydophila pneumoniac J138]	8.3	
		gi 12382001 dbj BAB21270.1  (AP002844) hypothetical		
3337	12382001	protein [Oryza sativa]	2.6	
		gi 11466493 ref[NP_038196.1  cytochrome c oxidase		
		subunit 3 [Chrysodidymus synuroideus]		
		gb AAF36962.1 AF222718_36 (AF222718) cytochrome		
3338	11466493	c oxidase subunit 3 [Chrysodidymus synuroideus]	5.4	
		gi 7475952 pir  H69754 two-component sensor histidine		
		kinase homolog yccG - Bacillus subtilis		
	ŀ	dbj BAA22234.1  (AB000617) YccG [Bacillus subtilis]		
		emb CAB12067.1  (Z99105) similar to two-component		
3339	7475952	sensor histidine kinase [YccH] [Bacillus subtilis]	5.9	
	ŀ	gi 340613 gb AAA31881.1  (L07545) A 'c' was inserted		
l .		after nt 369 (=nt 10459 in genomic sequence (M10126))		
		to correct -1 frameshift probably due to gel compression		
3344	340613	[Leishmania tarentolae]	1.4	
3345	10121788	gi 10121788 gb AAG13373.1  (AF268180) polyprotein [bovine viral diarrhea virus type 2]	- 4	
3343	10121788	[bovine virai diarrnea virus type 2]	5.4	
		ail2641252 ab14 A C26218 11 (A F052127) t		
3346	3641252	gi 3641252 gb AAC36318.1  (AF053127) leucine-rich receptor-like protein kinase [Malus x domestica]	0.49	
3340	3041232		0.49	
1		gi 6226170 sp O67271 SYE_AQUAE GLUTAMYL-		
1		TRNA SYNTHETASE (GLUTAMATE-TRNA		
		LIGASE) (GLURS) pir  D70405 glutamatetRNA ligase (EC 6.1.1.17) - Aquifex aeolicus		
		gb AAC07230.1  (AE000729) glutamyl-tRNA		
3347	6226170	synthetase [Aquifex aeolicus]	3	
3377	0220170	gi 2127912 pir  B64428 formate hydrogenlyase, subunit		
	1	gi 212/912 pir  B64428 formate hydrogenlyase, subunit   5 - Methanococcus jannaschii gb AAB99031.1		
	1	3 - Methanococcus Jannaschii gb AAB99031.1    (U67545) formate hydrogenlyase, subunit 5		
3348	2127012	[Methanococcus jannaschii]	2.9	
3340	212/712	Listomanococcus jannascinij	4.7	

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#### Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins) SEO ID ACCESS NO N DESCRIPTION P VALUE gi|7507391|pir||T24665 hypothetical protein T07F10.4 -Caenorhabditis elegans emblCAB01241.11 (Z77669) predicted using Genefinder-Similarity to Human mRNA 3351 7507391 (KIAA0033) (TR:G436224) [Caenorhabditis elegans] 6.3 gi|7476137|pir||A70989 hypothetical glycine-rich protein Rv1768 - Mycobacterium tuberculosis (strain H37RV) emblCAB09311.11 (Z95890) PE PGRS 3355 7476137 [Mycobacterium tuberculosis] 4.1 gi|6782280|emb|CAB70102.1| (AL132949) Y53F4B.7 [Caenorhabditis elegans] emb[CAB70113.2] 6782280 (AL132949) Y53F4B.26 [Cacnorhabditis elegans] 3359 5.6 gi|14133241|dbj|BAA86438.2| (AB032950) KIAA1124 3361 14133241 protein [Homo sapiens] 0.28 gil117000lsplP04371lCOX1 TRYBB CYTOCHROME C OXIDASE POLYPEPTIDE I pirlODUTMB cytochrome-c oxidase (EC 1.9.3.1) chain I -Trypanosoma brucei mitochondrion gb|AAB59223.1| (M14820) cytochrome c oxidase subunit I [Trypanosoma brucei] emb[CAB57806.1] (X01094) put. gene for cytochrome c oxidase subunit I gene 3362 117000 [Trypanosoma brucei] 1.7 gi|2072972|gb|AAC51276.1| (U93572) putative p150 3369 2072972 [Homo sapiens] 0.000008 gi|13476092|ref|NP 107662.1| ABC transporter permease protein [Mesorhizobium loti] dbj|BAB53448.1| (AP003011) ABC transporter 3371 13476092 permease protein [Mesorhizobium loti] 0.43 gil13385538|reffNP 080316.1| RIKEN cDNA 2810036K01 gene [Mus musculus] dbi|BAB28520.1] 3372 13385538 (AK012865) putative [Mus musculus] 2E-58 gi|14765434|ref|XP 030354.1| similar to KIAA0220 3376 14765434 protein (H. sapiens) [Homo sapiens] 8 gi|10641134|dbi|BAB16313.1| (AB049587) fork 3393 10641134 head/HNF-3 homologue [Ciona savignyi] 9.2 gi|11358605|pir||T49882 pectin methyl-esterase-like protein - Arabidopsis thaliana emb|CAB87932.1| (AL163912) pectin methyl-esterase-like protein 3396 11358605 [Arabidopsis thaliana] 4.1

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	, and the same	,
NO	N	DESCRIPTION	P VALUE
		gi 4493932 emb CAB38968,1  (AL034556)	
	İ	Hypothetical protein, PFC0580c [Plasmodium	
3397	4493932	falciparum]	7
		gi 11496667 ref NP 045466.1  conserved hypothetical	
	Ì	protein [Borrelia burgdorferi] pirlF70231 conserved	
		hypothetical protein BBG06 - Lyme disease spirochete	
		plasmid G/lp28-2 gb AAC66054.1  (AE000786)	
3400	11496667		7.2
3400	11490007	gi 7020787 dbi BAA91274.1  (AK000589) unnamed	7.3
3403	7020787	protein product [Homo sapiens]	6T 10
3403	7020787		5E-19
		gi 12045115 ref NP_072926.1  lipoprotein, putative	
		[Mycoplasma genitalium] sp P47502 Y260_MYCGE	
		HYPOTHETICAL LIPOPROTEIN MG260	
		PRECURSOR pir   G64228 hypothetical protein	
		homolog MG260 - Mycoplasma genitalium	
		gb AAC71481.1  (U39705) lipoprotein, putative	
3404	12045115	[Mycoplasma genitalium]	3.4
		gi 13959004 gb AAK51055.1 AF361075_2 (AF361075)	
3407	13959004	UL24 [Canine herpesvirus]	0.89
		gi 13814730 gb AAK41724.1  (AE006765) Hypothetical	
3408	13814730	protein [Sulfolobus solfataricus]	9.1
		gi 12744896 gb AAK06840.1 AF329637_1 (AF329637)	
3415	12744896	mitofusin 1 precursor [Homo sapiens]	0.16
		gi 8922500 ref NP_060600.1  hypothetical protein	
		FLJ10539 [Homo sapiens] dbj BAA91669.1	
3416	8922500	(AK001401) unnamed protein product [Homo sapiens]	7.1
		gi 6635084 emb CAB64573.1  (AL135930) hypothetical	
3418	6635084	protein L4738.02 [Leishmania major]	3.5
		gi 296559 cmb CAA49925.1  (X70529) ORF YBR1730	
3419	296559	[Saccharomyces cerevisiae]	0.42
		gi 5579432 gb AAD45553.1 U70376_18 (U70376)	
3422	5579432	SpcH [Streptomyces netropsis]	6.3
		gi 11466493 refINP 038196.1  cytochrome c oxidase	
		subunit 3 [Chrysodidymus synuroideus]	
		gb AAF36962.1 AF222718 36 (AF222718) cytochrome	
3423	11466493	c oxidase subunit 3 [Chrysodidymus synuroideus]	6
		gi 1084985 pir  S51908 cryptogene protein G1(ND9) -	
3424	1084985	Leishmania tarentolae (strain LEM125)	6.6
		gi 8778413 gb AAF79421.1 AC025808 3 (AC025808)	
3426	8778413	F18O14.6 [Arabidopsis thaliana]	4.2
J .20	2770113	[ Additiopsis triumina]	7.2

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#### Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins) SEO ID ACCESS NO N DESCRIPTION P VALUE gi|13992688|gb|AAK51582.1|AC022352 18 3430 13992688 (AC022352) Putative retroelement [Oryza satival] 9.3 gi|9280694|gb|AAF86563.1|AC069252 22 (AC069252) 3431 9280694 F2E2.6 [Arabidopsis thaliana] 3.3 gi|4115918|gb|AAD03429.1| (AF118222) similar to nascent polypeptide associated complex alpha chain 3437 4115918 [Arabidopsis thaliana] 4.4 gi|7433244|pir||T01751 gibberellin 20-oxidase - common tobacco dbj|BAA31690.1| (AB016084) Ntc16 3438 7433244 [Nicotiana tabacum] 1.2 gi|3978489|gb|AAC83366,1| (AF092918) unknown 3439 3978489 [Pseudomonas alcaligenes] 8.5 gi|7506359|pir||T23969 hypothetical protein R06C7.5 -Caenorhabditis elegans emb|CAA95843.1| (Z71266) Similarity to Human adenylosuccinate lyase (SW:PUR8 HUMAN), contains similarity to Pfam domain: PF00206 (Lvase), Score=165.2, E-value=3.5e-46, N=1~cDNA EST EMBL:Z14522 comes from this gene~cDNA EST yk67g5.3 comes from this 7506359 gene~cDNA EST yk67g5> 3441 5.9 gi|12852662|dbj|BAB29494.1| (AK014667) putative 3443 12852662 [Mus musculus] 2E-28 gi|9256527|ref[NP 061764.1| ceroid-lipofuscinosis. neuronal 8 (epilepsy, progressive with mental retardation) [Homo sapiens] gb|AAF13115,1|AF123757 1 (AF123757) putative transmembrane protein [Homo sapiens] gblAAF13116.1lAF123758 1 (AF123758) putative transmembrane protein [Homo sapiens] gb|AAF13117.1|AF123759 1 (AF123759) putative transmembrane protein [Homo sapiens] gb|AAF13118.1|AF123760 1 (AF123760) putative transmembrane protein [Homo saniens] gb|AAF13119.1|AF123761 1 (AF123761) putative 3448 9256527 transmembrane protein [Homo sapiens] 2.7 gi|7705167|gb|AAC60545,2| (S54379) sucrose-3457 7705167 phosphate synthase; SPS [Spinacia oleracea] 8.6 gi|10179322|dbi|BAB13673.1| (AB041350) type IV 3461 10179322 collagen alpha 5 chain [Mus musculus] 0.25 gi|13882453|gb|AAK47018.1| (AE007102) hypothetical 3462 13882453 protein [Mycobacterium tuberculosis CDC1551] 36

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
-110		DESCRIPTION	PVALUE
İ	}	gi 2499996 sp Q45298 PTGA CORGL PTS SYSTEM	
	1	GLUCOSE-SPECIFIC IIABC COMPONENT	
ļ	ł	(EIIABC-GLC) (GLUCOSE-PERMEASE IIABC	
i	İ	COMPONENT) (PHOSPHOTRANSFERASE	
		ENZYME II, ABC COMPONENT) (EII-GLC/EIII-	
		GLC) gb AAA22992.1  (L18875) phosphoenolpyruvate	í
		sugar phosphotransferase [Corynebacterium	'
3466	2499996	glutamicuml	9,9
	2,,,,,,	gi 7503384 pir  T31714 probable zinc proteinase (EC	5.5
	1	3.4.24) F44E7.4 - Caenorhabditis elegans	
		gb[AAC25789.1] (AF016421) Hypothetical protein	
3467	7503384	F44E7.4 [Caenorhabditis elegans]	5.1
3407	7505564	gi 225485 prf  1304284B beejin [Dictyostelium	3.1
3472	225485	discoideum]	5
3412	223403	gi 7302644 gb AAF57724.1  (AE003800) fj gene	
3482	7302644	product [Drosophila melanogaster]	0.19
3402	7302044	gil3121914 splO37679 COX3 THEAN	0.19
		CYTOCHROME C OXIDASE POLYPEPTIDE III	
ĺ	1	gb AAA73631.1  (U32225) cytochrome oxidase	
3487	3121914		6.9
5107	3121714	pospopulae in [Theneria annuala.]	0.7
	1	gi 1351243 sp P47749 THRR XENLA THROMBIN	
		RECEPTOR PRECURSOR pir  151667 thrombin	
	1	receptor - African clawed frog gb AAA18498.1	
3489	1351243	(U09632) thrombin receptor [Xenopus laevis]	6.2
		gi 12517831 gb AAG58344.1 AE005549 1 (AE005549)	
		aerobic respiration sensor-response protein; histidine	
	1	protein kinase/phosphatase, sensor for arcA	
		[Escherichia coli O157:H7 EDL933] dbj[BAB37512.1]	
	1	(AP002564) aerobic respiration sensor-response protein	
3491	12517831	[Escherichia coli O157:H7]	6,9
		gi 2240037 gb AAB66910.1  (AF005356) integrin	
3492	2240037	subunit betaCn1 [Acropora millepora]	5.5
3494	14742385	gi 14742385 ref XP_046349.1  33219 [Homo sapiens]	4.6
		gi 12847027 dbj BAB27408.1  (AK011113) putative	
3501	12847027	[Mus musculus]	5.6
		gi 1763691 gb AAB39835.1  (U72048) glycoprotein	
3505	1763691	[Classical swine fever virus]	1.2
		gi 13470696 ref NP_102265.1  unknown protein	
		[Mesorhizobium loti] dbj BAB48051.1  (AP002994)	
3507	13470696	unknown protein [Mesorhizobium loti]	5.1

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS	DESCRIPTION	
NU	N	DESCRIPTION	P VALUE
		gi 7706335 ref NP_057140.1  CGI-119 protein [Homo	
3511	7706335	sapiens] gb AAD34114.1 AF151877_1 (AF151877)	
3311	7706335	CGI-119 protein [Homo sapiens]	2.3
		gi 13812236 ref NP_113367.1  hypothetical protein	
2512	1201222	[Guillardia theta] gb AAK39923.1 AF165818_131	
3512	13812230	(AF165818) hypothetical protein [Guillardia theta]	2.4
		  gi 7510320 pir  T33571 hypothetical protein Y59C2A.2 -	
		Caenorhabditis elegans gb AAC68742.1  (AF099003)	
3513	7510220	Y59C2A.2 gene product [Cacnorhabditis elegans]	0.37
3313	7310320	gi 12724149 gb AAK05281,1 AE006350 8 (AE006350)	0.37
		HYPOTHETICAL PROTEIN [Lactococcus lactis	
3517	12724140	subsp. lactis]	1.7
3317	12/24149	gi 2342679 gb AAB70402.1  (AC000106) Similar to	1.7
		Vicia sativa ENBP1 (gb X95995). [Arabidopsis	
3519	2342679	thalianal	5.2
3317	2342019	gi 7305389 ref NP 038658.1  polycystic kidney disease	3,2
		1 homolog; polycystin-1 [Mus musculus]	
		gb[AAC53207.1] (U70209) polycystic kidney disease 1	
3520	7305389	protein [Mus musculus]	2.2
3320	7303369	gi[1353257]gb[AAB06234.1] (U26665) dimethyl	2.2
		sulphoxide reductase subunit B [Haemophilus	
3523	1353257	influenzae]	3,9
3323	1555257	mmucizacj	3,5
		gi 14794474 gb AAK73355.1 AF390546 1 (AF390546)	
3524	14794474	gut-enriched kruppel-like factor [Rattus norvegicus]	0.47
		See everyon method frames not egrend	
		gi 7510320 pir  T33571 hypothetical protein Y59C2A.2 -	
		Caenorhabditis elegans gb AAC68742.1  (AF099003)	
3527	7510320	Y59C2A.2 gene product [Caenorhabditis elegans]	0.89
		gi 10173181 dbj BAB04286.1  (AP001509) nickel	
		transport system (nickel-binding protein) [Bacillus	
3530	10173181	halodurans]	4.6
			- "-
		gi 7442453 pir  G72215 oligopeptide ABC transporter,	
]		permease protein - Thermotoga maritima (strain MSB8)	
		gb AAD36813.1 AE001813 5 (AE001813) oligopeptide	
		ABC transporter, permease protein [Thermotoga	
3531	7442453	maritima]	8.7
1		gil11136027 sp O88553 ZF37 RAT ZINC FINGER	
		PROTEIN 37 (ZFP-37) gb AAC24590.1  (AF072439)	
3532	11136027	zinc-finger protein-37; ZFP-37 [Rattus norvegicus]	1.8

	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESS				
NO	N	DESCRIPTION	P VALUE		
1		gi 14732500 ref XP_041698.1  hypothetical protein			
		FLJ10904 [Homo sapiens] ref[XP_041699.1  50659			
3538	14732500	[Homo sapiens]	8.5		
		gi 7438506 pir  T09963 mitosis-specific cyclin B-type -			
		Madagascar periwinkle dbj BAA20411.1  (D86386) B-			
3540	7438506	type cyclin [Catharanthus roseus]	0.75		
		gi 7495199 pir  T31857 hypothetical protein C02E7.14 -			
		Caenorhabditis elegans gb AAC24171.1  (AF016446)			
		Hypothetical protein C02E7.14 [Caenorhabditis			
3541	7495199	elegans]	4.9		
		gi 162223 gb AAA30233.1  (M27163) RNA polymerase			
3542	162223	III [Trypanosoma brucei]	4.9		
		gi 6635084 cmb CAB64573.1  (AL135930) hypothetical			
3546	6635084	protein L4738.02 [Leishmania major]	2.7		
		gi 13095647 ref NP_076562.1  unknown [Bovine			
		herpesvirus 4] gb AAK07989.1 AF318573_69			
3549	13095647	(AF318573) unknown [Bovine herpesvirus 4]	6.9		
i		gi 11359423 pir  T48729 hypothetical protein 8D4.30			
		[imported] - Neurospora crassa emb CAB88545.1			
		(AL353819) conserved hypothetical protein			
3551	11359423	[Neurospora crassa]	5.3		
		gi 13241881 gb AAK16450.1  (AF327992) cytochrome			
3556	13241881	oxidase subunit I [Platycryptus undatus]	9.7		
		gi 4589564 dbj BAA76804.1  (AB023177) KIAA0960			
3562	4589564	· · · · · · · · · · · · · · · · · · ·	4.8		
		gi 7486330 pir  T05113 hypothetical protein			
		F28M20.240 - Arabidopsis thaliana emb CAA19766.1			
		(AL031004) putative protein [Arabidopsis thaliana]			
2562	740/220	emb CAB79875.1  (AL161579) putative protein			
3563	7486330	[Arabidopsis thaliana]	8.3		
		gi 118965 sp P23098 DYHC_TRIGR DYNEIN BETA			
		CHAIN, CILIARY pir S17653 dynein beta heavy			
		chain, ciliary - sea urchin (Tripneustes gratilla)			
		cmb CAA42170.1  (X59603) Beta heavy chain of outer- arm axonemal dynein ATPase [Tripneustes gratilla]			
		prf  1714372A dynein:SUBUNIT=beta heavy chain			
3569	118965	Tripneustes gratillal	7.7		
3303	110703	gi 7662078 ref NP 055454.1  KIAA0372 gene product	1.1		
		[Homo sapiens] dbi[BAA20827,1] (AB002370)			
3571	7662078		5E-65		
3311	.002076	rent at 200 . 2. [violito subjected]	JE-03		

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 5803098 ref NP_006757.1  zinc finger protein 220;	
	ļ.	Monocytic leukemia zinc finger protein [Homo sapiens]	
		sp Q92794 MOZ_HUMAN MONOCYTIC	
		LEUKEMIA ZINC FINGER PROTEIN (ZINC	
		FINGER PROTEIN 220) gb AAC50662.1  (U47742)	
3576	5803098	monocytic leukaemia zinc finger protein [Homo sapiens]	4.6
	1	gi 9964384 ref NP_064852.1  AMV070 [Amsacta	
		moorei entomopoxvirus] gb AAG02776.1 AF250284_70	
		(AF250284) AMV070 [Amsacta moorei	
3582	9964384	entomopoxvirus]	9
	1		
0.500		gi 13272342 gb AAK17105.1 AF291051_2 (AF291051)	1.0
3583	13272342	ATP synthase subunit a [Candidatus Carsonella ruddii]	1.3
		gi 7505649 pir  T28783 hypothetical protein K09E10.1 -	
		Caenorhabditis elegans gb AAC48044.1  (AF003149)	
3586	7505649	Hypothetical protein K09E10.1 [Caenorhabditis elegans]	2.9
3360	7303649	gi 2981631 dbj BAA25253.1  (AB012223) ORF2 [Canis	2.9
3587	2001621	familiaris] (AB012223) OKF2 [Callis	0.98
3367	2961031	gi 6018210 gb AAF01791.1  (AF143853) 82 kD heat	0.56
3590	6018210	shock protein [Moniliformis moniliformis]	0.15
3370	0010210	shock protom [wommornus mountornus]	0.10
		gi 7160719 emb CAA88053,2  (Z48007) contains	
		similarity to Pfam domain: PF00069 (Eukaryotic protein	
	ì	kinase domain), Score=49.7, E-value=3.4e-13, N=3;	
		PF00211 (Adenylate and Guanylate cyclase catalytic	
		domain), Score=325.1, E-value=2.5e-94, N=1;	
3593	7160719	PF01094 (Receptor family ligand bi>	4.3
		gi 7470593 pir  S75491 hypothetical protein slr2115 -	
		Synechocystis sp. (strain PCC 6803) dbj[BAA18052.1]	
		(D90911) ORF_ID:slr2115~unknown protein	
3599	7470593	[Synechocystis sp. PCC 6803]	5.5
		gi 1334398 emb CAA33190.1  (X15081) MURF2	
3600	1334398	protein (AA 1-348) [Crithidia fasciculata]	2.2

[		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	Treatest Tresgators (Stable 13, 110); Redithum	Trotonis
NO	N	DESCRIPTION	P VALUE
		gi 118965 sp P23098 DYHC_TRIGR DYNEIN BETA	
ŀ	ľ	CHAIN, CILIARY pir  S17653 dynein beta heavy chain, ciliary - sea urchin (Tripneustes gratilla)	
	,	emb CAA42170.1  (X59603) Beta heavy chain of outer-	
		arm axonemal dynein ATPase [Tripncustes gratilla] prf  1714372A dynein:SUBUNIT=beta heavy chain	
3602	118965	[Tripncustes gratilla]	9.8
		gi 14578630 gb AAK68919,1 AF189282 9 (AF189282)	
3603	14578630	putative glycosyltransferase [Bacteroides fragilis]	0.29
		gi 14318569 ref NP_116702.1  Yfr044cp [Saccharomyces cerevisiae] sp P43616 YFL4 YEAST	
		HYPOTHETICAL 52.9 KD PROTEIN IN SAP155-	
		YMR31 INTERGENIC REGION pir S56299	
		hypothetical protein YFR044c - yeast (Saccharomyces	
		cerevisiae) dbj BAA09283.1  (D50617) YFR044C	
3606	14318569	[Saccharomyces cerevisiae]	33
3609	10581847	gi 10581847 gb AAG20525.1  (AE005124) Vng2444c [Halobacterium sp. NRC-1]	4,5
		gi 1363925 pir  S57662 hypothetical protein 2 - North	
3616	1363925	American opossum (fragment) emb CAA88817.1  (Z48955) ORF-2, putative RT [Didelphis virginiana]	3.6
		gi 1808609 cmb CAA64091.1  (X94355) D6L [Cowpox	
3618	1808609	virus] emb CAA72556.1  (Y11842) C6L [Cowpox virus]	0.6
3018	1000009	gi[14775653]ref[XP_045559.1] similar to KIAA0565	9.6
3619	14775653	gene product (H. sapiens) [Homo sapiens]	4.7
		gi 7475003 pir  G69801 hypothetical protein yfhO - Bacillus subtilis emb CAB12689.1  (Z99108) yfhO	
		Bacillus subtilis dibjBAA24481,1 (D85082) YfhO	
3621	7475003	[Bacillus subtilis]	8.1
		gi 5902895 dbj BAA84478.1  (AB032367) type I	
3626	5902895	polyketide synthase AVES 3 [Streptomyces avermitilis]	4.3
1		gi[7444049 pir  F72275 phosphate regulon transcription	
		regulator PhoB - Thermotoga maritima (strain MSB8) gb AAD36333.1 AE001781 4 (AE001781) phosphate	
		regulon transcriptional regulatory protein PhoB	
3627	7444049	[Thermotoga maritima]	2.8

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundam	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
	Ì	gi 7507776 pir  T16867 probable cytochrome P450	
3630	7507776	T13C5.1 [similarity] - Caenorhabditis elegans	1.6
	1		
	ļ	gi 11360480 pir  H82745 acyl-[ACP]-UDP-N-	[
	1	acetylglucosamine XF0918 [imported] - Xylella	
	l	fastidiosa (strain 9a5c) gb AAF83728.1 AE003931_5	
		(AE003931) acyl-[ACP]-UDP-N-acetylglucosamine	
3632	11360480	[Xylella fastidiosa 9a5c]	1,3
		gi 13873167 gb AAK43406.1  (AF196891)	
		polygalacturonase inhibitor protein [Fragaria vesca]	
		gb AAK43407.1  (AF196892) polygalacturonase	
		inhibitor protein [Fragaria vesca] gb AAK43408.1	
		(AF196893) polygalacturonase inhibitor protein	
3633	13873167	[Fragaria vesca]	3.7
		gi 10946710 ref NP_067350.1  Rhesus blood group-	
		associated B glycoprotein; Rh type B glycoprotein [Mus	
		musculus] gb AAF19371.1  (AF193808) Rh type B	
3636	10946710	glycoprotein [Mus musculus]	4.9
		gi 7447412 pir  T06201 xyloglucan endo-1,4-beta-D-	
		glucanase (EC 3.2.1) - barley emb CAA63662.1	
		(X93174) xyloglucan endotransglycosylase (XET)	
3638	7447412	[Hordeum vulgare]	5.5
		gi 12620601 gb AAG60877.1 AF322012_182	
3639	12620601	(AF322013) ID409 [Bradyrhizobium japonicum]	6.4
		gi 1175418 sp Q09743 ST16_SCHPO STE16	
		PROTEIN pir T39379 sexual differentiation and	
		meiosis protein ste20 - fission yeast	
		(Schizosaccharomyces pombe) emb CAA90815.1	
		(Z54140) required for sexual differentiation and meiosis	
		[Schizosaccharomyces pombe] emb CAA11758.1	
		(AJ223984) Ste20 protein [Schizosaccharomyces	
3640	1175418	pombe]	4.6
		gi 1172876 sp P35906 RDS_FELCA PERIPHERIN	
		(RETINAL DEGENERATION SLOW PROTEIN)	
	1170051	pir  I46087 peripherin - cat gb AAA19175.1  (M94047)	
3641	1172876	peripherin [Felis catus]	5,1
3640	00550	gi 825671 cmb CAA34809.1  (X16934) B23	
3643	825671	nucleophosmin (280 AA) [Homo sapiens]	2E-10
2545	20005#4	gi 2388576 gb AAB71457.1  (AC000098) YUP8H12.17	
3645	2388576	[Arabidopsis thaliana]	3.7

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
		gi 7498664 pir  T20598 hypothetical protein F08G2.4 -	
		Caenorhabditis elegans emb[CAB04058.1] (Z81495)	
		predicted using Genefinder~cDNA EST yk401a9.3	
ł		comes from this gene~cDNA EST yk401a9,5 comes	
		from this gene~cDNA EST yk523d8.3 comes from this	
		gene~cDNA EST yk523d8.5 comes from this gene	
		[Caenorhabditis elegans] gb AAG50226.1 AF303268_1	
3648		(AF303268) 2O123 [Cacnorhabditis elegans]	9.4
3654	225992	gi 225992 prf  1405326A GLI gene [Homo sapiens]	3.2
		gi 422532 pir  A45407 collagen alpha 3(IV) chain - sea	
3657	422532	urchin (Strongylocentrotus purpuratus)	0.94
		gi]14149807 ref[NP_115517.1  hypothetical protein	
		DKFZp434K1421 [Homo sapiens] emb CAB66740.1	
3658	14149807	(AL136806) hypothetical protein [Homo sapiens]	4E-65
		gi 134872 sp P13666 SRCA_RABIT	
		SARCALUMENIN PRECURSOR gb AAA31189.1	
		(M25750) sarcolumenin precursor [Oryctolagus	
3665	134872	cuniculus]	8.1
2670	14504055	2145240551 SYP 046550 114 1 FF	0.45
3670	14734955	gi]14734955 ref[XP_046758.1  tensin [Homo sapiens]	0.47
3675	14742385	gi 14742385 ref XP 046349.1  33219 [Homo sapiens]	2.1
		gi 2197085 gb AAD04635.1  (AF003535) ORF2-like	
3676	2197085	protein [Homo sapiens]	0.14
		gi 1711034 gb AAB38323.1  (U78953) basic helix-loop-	
		helix DNA binding protein HLH-3 [Caenorhabditis	
3690	1711034	clegans]	5.4
		gi 2072964 gb AAC51271.1  (U93569) putative p150	
3692	2072964	[Homo sapiens]	0.15
		gi 1857258 gb AAB48409,1  (U75538) putative viral	•
3693	1857258	polymerase [tobacco streak virus]	5.3
		gi 8573079 ref NP_059567.1  NADH dehydrogenase	
		subunit 1 [Sciurus vulgaris] emb CAB93985.1	
		(AJ238588) NADH dehdrogenase subunit 1 [Sciurus	
3694	8573079	vulgaris]	3E-15
		gi 12654875 gb AAH01281.1 AAH01281 (BC001281)	
l		tumor necrosis factor receptor superfamily, member 10b	
3699	12654875	[Homo sapiens]	7.9
2001		gi 2961586 gb AAC05758.1  (AF051294) NADH	
3704	2961586	dehydrogenase subunit 1 [Dalbulus gelbus]	4.3

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	,	
NO	N	DESCRIPTION	P VALUE
	l		
2505	1504016	gi 1504016 dbj BAA13207.1  (D86971) no similarities	
3707	1504016	to reported gene products [Homo sapiens]	6E-11
	Į.	gi 11466493 ref[NP_038196.1  cytochrome c oxidase	
		subunit 3 [Chrysodidymus synuroideus]	
3709	11466402	gb AAF36962.1 AF222718_36 (AF222718) cytochrome c oxidase subunit 3 [Chrysodidymus symroideus]	
3709	11400493	gi 14089788 cmb CAC13547.1  (AL445564)	5.7
3710	14000700	LIPOPROTEIN [Mycoplasma pulmonis]	2.4
3/10	14009700	gi 8978260 dbi BAA98151.1  (AB025612) CLP	2.4
		protease regulatory subunit CLPX-like [Arabidopsis	
3724	8978260	thalianal	5.4
3124	07/0200	gi 14748431 ref XP 018068.2  hypothetical protein	3,4
3727	14748431	MGC3199 [Homo sapiens]	0,000009
3121	14740431	gi 10727054 gb AAF46943.2  (AE003459) CG3037	0.000009
3734	10727054	gene product [Drosophila melanogaster]	3.8
3731	10727054	gene product [Drosopinia motatogaster]	5.0
	1	gi 6323629 ref NP 013700.1  Yml013c-ap	
		[Saccharomyces cerevisiae] pir S69871 hypothetical	
3735	6323629	protein YML012c-a - yeast (Saccharomyces cerevisiae)	9
	0020027	gi 4502081 ref[NP 001626,1] amphiphysin;	
		amphiphysin (Stiff-Man syndrome with breast cancer	
		128kD autoantigen) [Homo sapiens] refXP 035768.1	
		amphiphysin [Homo sapiens] ref[XP_035768.1]	
		amphiphysin [Homo sapiens]	
		spiP49418iAMPH HUMAN AMPHIPHYSIN	
	ļ	pir  S62400 amphiphysin (clone 22-2) - human	
		gb AAA21865.1  (U07616) amphiphysin [Homo	
		sapiens] emb[CAA57197.1] (X81438) amphiphysin	
3736	4502081	[Homo sapiens]	0.61
		gi 7492913 pir T40695 probable ribosomal protein	
	l	precursor, mitochondrial - fission yeast	
		(Schizosaccharomyces pombe) emb CAB36868.1	
		(AL035536) mitochondrial ribosomal protein L36	
3741	7492913	[Schizosaccharomyces pombe]	6.7
		gi 1168945 sp P30572 CHS2_CANAL CHITIN	
		SYNTHASE 2 (CHITIN-UDP ACETYL-	
		GLUCOSAMINYL TRANSFERASE 2)	
		gb AAB59308.2  (M82937) chitin synthase 2 [Candida	
3743	1168945	albicans]	1.2

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Donate la ch
SEQ ID	ACCESS	Table 3B Realest Reignbor (BlastA vs. Non-Redundam	Proteins)
NO	N	DESCRIPTION	P VALUE
		gi 12722927 gb AAK04181.1 AE006247 3 (AE006247)	
		UNKNOWN PROTEIN [Lactococcus lactis subsp.	
3745	12722927	lactis]	2.9
		gi 6090792 gb AAF03328.1  (AF101747) olfactory	
3747	6090792	receptor [Pan paniscus]	4.1
		gi 12846356 dbj BAB27137.1  (AK010713) putative	
3750	12846356	[Mus musculus]	0.000000004
		gi 4240279 dbj BAA74918.1  (AB020702) KIAA0895	
3754	4240279	protein [Homo sapiens]	10
		gi 543715 sp P36875 2AAA_PEA PROTEIN	
1		PHOSPHATASE PP2A REGULATORY SUBUNIT A	
		(PR65) pir  S40171 phosphoprotein phosphatase 2A	
1		65kDa regulatory chain - garden pea (fragment)	
		pir  S43776 phosphoprotein phosphatase 2A 65kDa	
		regulatory subunit - garden pea (fragment)	
		emb CAA81107.1  (Z25888) phosphoprotein	
		phosphatase 2A 65kDa regulatory subunit [Pisum	
3755	543715	sativum]	6E-12
Ì			
		gi 11353649 pir  D81094 hypothetical protein NMB1340	
	1	[imported] - Neisseria meningitidis (group B strain	
2000	11252640	MD58) gb AAF41715.1  (AE002482) hypothetical	
3756	11353649	protein [Neisseria meningitidis MC58]	3.4
Ì		gi 11276053 gb AAG33872.1 AF319939_1 (AF319939)	
3761	11276053		4.3
3701	11270033	cinalen [enoristolicula fullificiana grandiovirus]	4.3
l		gi 13812312 ref[NP 113430.1  40S ribosomal protein	
	i .	SSA [Guillardia theta] emb CAC26999.1  (AJ010592)	
3763	13812312	40S ribosomal protein SSA [Guillardia theta]	9,3
2703	13012512	gi 4115943 gb AAD03453.1  (AF118223) contains	
		similarity to eukaryotic protein kinase domains (Pfam:	
		PF00069, score=312.6, E=4.7e-90, N=1) and EF hand	
	\	domains (Pfam: PF00036, score=131, E=2.1e-35, N=4)	
		[Arabidopsis thaliana] emb[CAB80837.1] (AL161501)	
i		putative calcium dependent protein kinase [Arabidopsis	
3764	4115943	thaliana]	1.2
		gi 7494381 pir  H71613 probable multiple	
		transmembrane domain protein PFB0475c - malaria	
		parasite (Plasmodium falciparum) gb AAC71884.1	
		(AE001397) predicted multiple transmembrane domain	
3765	7494381	protein [Plasmodium falciparum]	8.5

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Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS	Tions of Tions of Tours of The Teaching	l Touchis)
NO	N	DESCRIPTION	P VALUE
			- 11202
	1	gi 3930075 gb AAC82262.1  (U81928) envelope	
3768	3930075	glycoprotein [Human immunodeficiency virus type 1]	2,5
		gi 4493896 cmb CAB39005.1  (AL034558) predicted	
		using hexExon; MAL3P2.18 (PFC0245c), Hypothetical	
3769	4493896	protein, len: 3934 aa [Plasmodium falciparum]	9.2
		gi 14193274 gb AAK55866.1 AF267203_2 (AF267203)	
		ATP synthase gamma subunit [Candidatus Carsonella	
3770	14193274	ruddii]	6.3
		gi 11499055 ref NP_070289.1  hypothetical protein	
		[Archaeoglobus fulgidus] sp O28812 YE60_ARCFU	
	1	HYPOTHETICAL PROTEIN AF1460 pir C69432	
		hypothetical protein AF1460 - Archaeoglobus fulgidus	
		gb AAB89791.1  (AE001002) A. fulgidus predicted	
3771	11499055	coding region AF1460 [Archaeoglobus fulgidus]	5.9
		gi 7301453 gb AAF56578.1  (AE003755) CG12290	
3773	7301453	gene product [Drosophila melanogaster]	9.3
		gi 11467606 ref NP_050068.1  NH2 terminus uncertain	
		[Leishmania tarentolae] pir  B26696 hypothetical protein	
		1 (CYb-COII intergenic region) - Leishmania tarentolae	
		mitochondrion (fragment) gb AAA96601.1  (M10126)	
3 <b>7</b> 77	11467606	NH2 terminus uncertain [Leishmania tarentolae]	0.4
		gi 7507710 pir  T23023 hypothetical protein T12A7.5 -	
		Caenorhabditis elegans emb CAB07423.1  (Z92847)	
		cDNA EST yk26a3.5 comes from this gene~cDNA EST	
		yk29g7.5 comes from this gene~cDNA EST yk29g8.5	
		comes from this gene~cDNA EST yk50c8.5 comes from	
		this gene~cDNA EST yk52f12.5 comes from this	
		gene~cDNA EST yk53c1.5 comes from this	
		gene~cDNA EST> emb CAA98143.1  (Z73911) cDNA	
		EST yk26a3.5 comes from this gene~cDNA EST	
		yk29g7.5 comes from this gene~cDNA EST yk29g8.5	
		comes from this gene~cDNA EST yk50c8.5 comes from this gene~cDNA EST yk52f12.5 comes from this	
		gene~cDNA EST yk52f12.5 comes from this	
3780	7507710	gene~cDNA EST>	2.6
5,00	.55,,20	Pare	4.0

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SEQ ID	ACCESS	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
NO NO	N N	DESCRIPTION	P VALUE
	1		
		gi 13994312 ref[NP_114137.1  testis transcript Y 13	
		[Homo sapicns] ref[XP_040549.1  testis transcript Y 13	
		[Homo sapiens] gb AAK13492.1 AF332242_1	
3783	13994312	(AF332242) transcript Y 13 [Homo sapiens]	2.6
		gi 1169093 sp P45815 CRF1_YARLI COPPER	
		RESISTANCE PROTEIN CRF1 cmb CAA80803.1	
		(Z23265) Product required for copper resistance	
3786	1169093		7.1
		gi 3599476 gb AAC69336.1  (AF084637) serendipity	
3787	3599476		2.5
_		gi 14728817 ref XP_047357.1  KIAA0342 gene product	
3793	14728817	[Homo sapiens]	9.5
		gi 11281706 pir  B81937 competence protein NMA0906	
		[imported] - Neisseria meningitidis (group A strain	
		Z2491) emb CAB84183.1  (AL162754) competence	
3795	11281706	protein [Neisseria meningitidis Z2491]	5.5
5,75	11201700	protein profesoria moningaidis 22491]	3.3
		gi 11096149 gb AAG30214.1 AF296334 1 (AF296334)	
3797	11096149	collagen-like surface protein [Streptococcus pyogenes]	2.4
2.77	11020112	gi 1827933 pdb 1PCA  Procarboxypentidase A	2.7
3798	1827933	(E.C.3.4:12.2)	1.3
		gi 6320364 ref NP_010444.1  regulator of transporters;	
		Ssy1p [Saccharomyces cerevisiae]	
		spiO03770ISSY1 YEAST PUTATIVE AMINO-ACID	
		PERMEASE SSY1 pir S57984 probable membrane	
		protein YDR160w - yeast (Saccharomyces cerevisiae)	
		emb CAA90380.1  (Z50046) unknown [Saccharomyces	
3800	6320364	cerevisiae]	0.92
		gi 14973521 gb AAK76078.1  (AE007490) ribosomal	÷
		large subunit pseudouridine synthase, RluD subfamily	
3803	14973521	[Streptococcus pneumoniae]	7.4
		gi 10834955 ref NP_066916.1  ICP4 protein [Gallid	
		herpesvirus 3] dbj BAB16594.1  (AB049735) ICP4	
3807	10834955	protein [Gallid herpesvirus 3]	1.6
		gi 13430085 gb AAK25738.1 AF291747_1 (AF291747)	
3817	13430085	zinc-finger transcription factor KROX20 [Gallus gallus]	0.63
		gi 7462422 pir  A72258 hypothetical protein -	
		Thermotoga maritima (strain MSB8)	
		gb AAD36479.1 AE001793_9 (AE001793) hypothetical	
3822	7462422		9.7

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESS				
NO	N	DESCRIPTION	P VALUE		
		gi 7517565 pir  A70328 hypothetical protein aq_311 -			
l		Aquifex aeolicus gb AAC06616.1  (AE000683) putative			
3828	7517565	protein [Aquifex acolicus]	3.7		
		gi 3924618 gb AAC79121.1  (U88902) putative			
3830	3924618	envelope protein [Homo sapiens]	0.0000006		
		gi 7023029 dbj BAA91807.1  (AK001647) unnamed			
3835	7023029	protein product [Homo sapiens]	4E-80		
		gi 12698001 dbj BAB21819.1  (AB051515) KIAA1728			
3836	12698001	protein [Homo sapiens]	0.0004		
		gi 5281354 gb AAD41491.1 AF136456_1 (AF136456)			
3837	5281354	transcription factor Tcf3b [Danio rerio]	5		
		gi 9628045 ref NP_042639.1  ORF 42 [Equine			
		herpesvirus 2] pir  S55637 hypothetical protein 42 -			
		equine herpesvirus 2 gb AAC13830.1  (U20824) ORF			
3838	9628045	42 [Equine herpesvirus 2]	7.1		
1 1		gi 9964462 ref NP_064930.1  AMV148 [Amsacta			
		moorei entomopoxvirus]			
2000	0061160	gb AAG02854.1 AF250284_148 (AF250284) AMV148			
3839	9964462	[Amsacta moorei entomopoxvirus]	5.8		
1 1					
		gi 13357611 ref NP_077885.1  membrane nuclease A -			
[ [		hypothetical [Ureaplasma urealyticum] pir  A82939			
		membrane nuclease A, hypothetical UU055 [imported] -			
1		Ureaplasma urealyticum gb AAF30460.1 AE002105_1			
3844	12257611	(AE002105) membrane nuclease A - hypothetical [Ureaplasma urealyticum]	20		
3644	13337611	gi 2425188 dbi BAA22281.1  (AB007035) FGF receptor	2.9		
3845	2425100	[3 [Xenopus lacvis]	2.5		
3043	2423100	3 [Actiopus facvis]	Z.3		
1		7112510501 : HTT02020			
[ [		gi 11351069 pir  H83232 probable ATP-dependent helicase PA3297 [imported] - Pseudomonas aeruginosa			
		(strain PAO1) gb AAG06685,1 AE004752 1			
1		(AE004752) probable ATP-dependent helicase			
3848	11351060	[Pseudomonas aeruginosa]	4.2		
20.0	11331007	gi 7141304 gb AAF37281.1  (AF225702) RSH1	1,2		
3850	7141304	[Arabidopsis thaliana]	9.3		
		[A MONGOPPE GARAGES			
1					
		gi 11352313 pir  G83376 probable trehalose synthase			
1 1		PA2152 [imported] - Pseudomonas aeruginosa (strain			
		PAO1) phIAAG05540 1IAE004642 7 (AE004642)			
3852	11352313	probable trehalose synthase [Pseudomonas aeruginosa]	3		
2952	11352313	PAO1) gb AAG05540.1 AE004642_7 (AE004642)	3		

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 9453863 dbj BAB03284.1  (AB037278) complement	
3856	9453863	C4A [Cyprinus carpio]	5,5
	l	gi 11466282 ref NP_049597.1  orf1386 [Tetrahymena	
	i	pyriformis] gb AAD41942.1 AF160864_30 (AF160864)	
3859	11466282	orf1386 [Tetrahymena pyriformis]	7.5
	l	gi 9626695 ref NP_040967.1  overlapping out-of-phase	ļ
		protein [Eggplant mosaic virus]	
		sp P20129 V70K_EPMV 70 KD PROTEIN	
		gb AAA43038.1  (J04374) overlapping out-of-phase	
3861	9626695	protein [Eggplant mosaic virus]	5.6
		gi 7473683 pir  H75456 probable succinate	
	1	dehydrogenase, hydrophobic subunit SdhD -	
	ĺ	Deinococcus radiodurans (strain R1)	
		gb AAF10526.1 AE001947_9 (AE001947) succinate	
		dehydrogenase, hydrophobic subunit SdhD, putative	
3862	7473683	[Deinococcus radiodurans]	5.2
	ļ	gi 11359953 pir  T46330 hypothetical protein	
	i	DKFZp434D0513.1 - human (fragment)	
		emb CAB70660.1  (AL137259) hypothetical protein	
3864	11359953	[Homo sapiens]	6E-28
		gi 3056600 gb AAC13911.1 AAC13911 (AC004255)	
3867	3056600	T1F9.21 [Arabidopsis thaliana]	5.5
		gi 11358605 pir  T49882 pectin methyl-esterase-like	
		protein - Arabidopsis thaliana emb CAB87932.1	
		(AL163912) pectin methyl-esterase-like protein	
3872	11358605	[Arabidopsis thaliana]	3.9
20.52	5015665	gi 5917666 gb AAD55980.1 AF159297_1 (AF159297)	
3873	5917666	extensin-like protein [Zea mays]	9.2
2070	15011400	gi 15011489 gb AAK77584.1 AF396436_24	
3879	15011489	(AF396436) heme maturase [Tetrahymena thermophila]	3
		gi 5835877 ref NP_008647.1 ND2_15072 NADH	
		dehydrogenase subunit 2 [Ceratitis capitata]	
2000	5025022	emb CAB45088.1  (AJ242872) NADH dehydrogenase	0.05
3880	5835877	(ubiquinone) chain 2 [Ceratitis capitata]	0.85

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	Table 5.5 Treat est Troignoot (Diasex vs. 1101-1101andant	1 Totalis)
NO	N	DESCRIPTION	P VALUE
		gi 10956089 ref NP 052581.1  Orf2 [Pediococcus	
		pentosaceus] ref[NP_037562.1] mobilization protein	
		[Pediococcus pentosaccus]	
		gb AAD25894.1 AF069302 2 (AF069302) Orf2	
		[Pediococcus pentosaceus]	
		gb AAD39619.1 AF033858 2 (AF033858) mobilization	
3881	10956089	protein [Pediococcus pentosaceus]	5.5
		gi 9294238 dbi BAB02140.1  (AP000411) contains	
		similarity to reverse transcriptase~gene id:K11J14.5	
3883	9294238	[Arabidopsis thaliana]	6.3
3003	7294230		0.3
		gi 975667 cmb CAA61500.1  (X89213) RNA	
		polymerase [Infectious hematopoietic necrosis virus]	
3885	975667	prf  2121413F RNA polymerase [Infectious	4.0
3003	9/3007	hematopoietic necrosis virus]	4.3
		gi 138293 sp P23041 VGLG HRSV8 MAJOR	
		SURFACE GLYCOPROTEIN G (ATTACHMENT	
		GLYCOPROTEIN G) pir  MGNZ60 major surface	
		glycoprotein G - human respiratory syncytial virus	
		(strain 8/60) gb AAA47408.1  (M73545) attachment	
		protein [Human respiratory syncytial virus]	
		gb AAA47413.1  (M55633) attachment glycoprotein	
3886	138293	[Human respiratory syncytial virus]	2.1
		gi 9628729 ref[NP 043760.1  U4 [Human herpesvirus 7]	
		sp P52521 VU4 HSV7J U4 PROTEIN pir T41908	
	l	hypothetical protein U4 - human herpesvirus 7 (strain	
		JI) gb[AAC54668.1] (U43400) U4 gene product [Human	
		herpesvirus 7] gb[AAC40722.1] (AF037218) U4	
3888	9628729	[Human herpesvirus 7]	0.77
		gi 13376093 ref NP_079033.1  hypothetical protein	
		FLJ12879 [Homo sapions] dbj BAB14321.1	
3891	13376093	(AK022941) unnamed protein product [Homo sapicns]	6.5
		gi 7508715 pir  T26003 hypothetical protein	
		VC27A7L.1 - Caenorhabditis elegans	
		emb[CAB09130.1] (Z95621) contains similarity to Pfam	
		domain: PF01461 (7TM chemoreceptor), Score=47.3,	
3892	7508715	E-value=1.9e-12, N=1 [Caenorhabditis elegans]	4.7
		gi 2961421 dbj BAA25153.1  (AB011027) Pns10-2 [oat	
3893	2961421	sterile dwarf virus]	7.5

		Table 2B Name + Night (Dig W N. B. J. J.	D +: \
CEO ID	ACCESS	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	N N	DESCRIPTION	DAZATITE
NO	- 1	DESCRIPTION	P VALUE
		gi 11276053 gb AAG33872.1 AF319939 1 (AF319939)	
3899	11276053	enhancin [Choristoneura fumiferana granulovirus]	3.7
3033	11270033	gi 10177870 dbi BAB11240.1  (AB010074)	3.7
	1	potassium/proton antiporter-like protein [Arabidopsis	
3904	10177870		2.7
	20111010		2.7
		gi 9629709 rcf NP 045001,1  methyltransferase (MT)	
		and helicase (HEL) domains [Little cherry closterovirus]	
*		emb CAA71285.1  (Y10237) methyltransferase (MT)	
3907	9629709	and helicase (HEL) domains [Little cherry closterovirus]	4.1
		gil14756677 reflXP 040326.1  absent in melanoma 1	
3908	14756677	[Homo sapiens]	0.96
		gi 9630708 ref[NP 047255,1  gag-pol precursor	
		polyprotein gPr80 [Feline leukemia virus]	
		gb AAC31801.1  (AF052723) gag-pol precursor	
3911	9630708	polyprotein gPr80 [Feline leukemia virus]	0.021
		gi 91039 pir  S04847 leukocyte adhesion protein CD18	
		precursor - mouse emb CAA33077.1  (X14951) CD18	
3913	91039	antigen preprotein [Mus musculus]	4.6
		gi 5107943 gb AAD40185.1 AF157488_1 (AF157488)	
		36DE accessory gland protein [Drosophila	
		melanogaster] gb AAF53664.1  (AE003658) Acp36DE	
3916	5107943	gene product [Drosophila melanogaster]	5.8
		gi 4809000 gb AAD30054.1  (AF133052) pol	
3919	4809000	polyprotein [walleye epidermal hyperplasia virus type 2]	0.000004
		gi 7262603 gb AAF43901.1 AF162134_1 (AF162134)	
3920	70/0/02	immunoglobulin epsilon heavy chain constant region [Felis catus]	8.9
3920	7202003		8.9
		gi 12963481 ref[NP_061927.1  hypothetical protein MGC5560; hypothetical protein [Homo sapiens]	
		dbj[BAB14342.1] (AK022978) unnamed protein product	
3927	12062481	[Homo sapiens]	2E-66
3921	12903401	[Fiolio sapiens]	ZE-00
3929	13651251	gi 13651251 ref XP_017240.1  15060 [Homo sapiens]	0.00000003
		gi 11466329 ref[NP 051157.1  ATP synthase F0 subunit	
		6 [Cafeteria roenbergensis]	
		gb AAF05808.1 AF193903_31 (AF193903) ATP	
3930	11466329	synthase F0 subunit 6 [Cafeteria roenbergensis]	8.3

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		Troums)
NO	N	DESCRIPTION	P VALUE
	1	gi 7452186 pir  S75936 hypothetical protein slr0023 -	
		Synechocystis sp. (strain PCC 6803) dbj BAA10783,1	
į	ļ	(D64006) ORF ID:slr0023~unknown protein	
3933	7452186		7.7
		gi 12720861 gb AAK02675,1  (AE006095) unknown	
3935	12720861	[Pasteurella multocida]	1.2
		gi 8919178 emb CAB96077.1  (AJ277244) alpha-	
3937	8919178	glucosidase [Solanum tuberosum subsp. tuberosum]	5.7
		gi 12830385 emb CAC29069.1  (AJ408289)	
		immunoglobulin lambda light chain variable region	
3938	12830385		2.1
		gi 13385576 ref[NP 080353.1  RIKEN cDNA	
	1	3110040N11 gene [Mus musculus] dbi BAB29184.1	
		(AK014163) putative [Mus musculus] dbj BAB31031.1	
	i	(AK018003) putative [Mus musculus] dbi BAB31634.1	
3942	13385576	(AK019261) putative [Mus musculus]	5E-48
		, , , , , , , , , , , , , , , , , , , ,	
		gi 13476785 ref[NP_108354.1  transcription regulator	
		[Mesorhizobium loti] dbi BAB53815.1  (AP003013)	
3945	13476785	transcription regulator [Mesorhizobium loti]	5,6
		gi 13357649 ref NP_077923.1  unique hypothetical	
		membrane lipoprotein [Ureaplasma urealyticum]	
		pir E82934 hypothetical protein UU092 [imported] -	
		Ureaplasma urealyticum gb AAF30498,1 AE002109 1	
		(AE002109) unique hypothetical membrane lipoprotein	
3947	13357649	[Ureaplasma urealyticum]	4.1
		[o-sq-same areas) areas	
		gi 13358144 ref NP 078418.1  conserved hypothetical	
		membrane lipoprotein [Ureaplasma urealyticum]	
[		pir  G82873 conserved hypothetical membrane	
		lipoprotein UU579 [imported] - Ureaplasma urealyticum	
		gb AAF30993.1 AE002156 8 (AE002156) conserved	
		hypothetical membrane lipoprotein [Ureaplasma	
3951	13358144	urealyticum]	8.1
		gi 7299437 gb AAF54626.1  (AE003691) KP78b gene	
3956	7299437	product [Drosophila melanogaster]	5.5
1		gi 6729023 gb AAF27019.1 AC009177 9 (AC009177)	
3961	6729023	hypothetical protein [Arabidopsis thaliana]	9,8

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESS				
NO	N	DESCRIPTION	P VALUE		
		gi 2645329 gb AAB87213.1  (U83821) NADH			
3962	2645329	dehydrogenase subunit 4 [Oryzomys palustris]	6.6		
		gi 11280919 pir  T46939 hypothetical protein 1			
		[imported] - Agrobacterium tumefaciens plasmid			
3965	11200010	pAtK84b gb AAD31598,1 AF065244_2 (AF065244)	0.22		
3903		unknown [Agrobacterium tumefaciens]	0.33		
		gi 7305249 ref NP_038748.1  MAX-interacting protein [Mus musculus] gb AAF24761.1  (AF205935) MGA			
3966	7305249	protein [Mus musculus]	3E-51		
3900	1303249	gi 5901753 gb AAD55397.1  (AF177905) hypothetical	312-31		
3969	5901753	protein [Candida glabrata]	1.6		
1 2200		gil9635492 ref[NP 059583.1  gtrA [Enterobacteria	***		
		phage P22] sp[P57021 GTRA BPP22			
		BACTOPRENOL-LINKED GLUCOSE			
		TRANSLOCASE gb AAF75001,1  (AF217253) gtrA			
3973	9635492	[Enterobacteria phage P22]	4		
		gi 2143428 pir  I58123 aggrecan - mouse (fragment)			
		gb AAB32159.1  (S73720) aggrecan=cmd(aggrecan)			
		[mice, cmd/cmd, liver, Peptide Partial Mutant, 112 aa]			
3975	2143428	[Mus sp.]	3.3		
		gi 14767565 ref XP_027089.1  N-acetylated alpha-			
		linked acidic dipeptidase 2 [Homo sapiens]			
		emb CAB39967.1  (AJ012370) NAALADase II protein			
3979	14767565	[Homo sapiens]	1.3		
		gi 11359267 pir  T50192 probable pseudouridylate			
		synthase [imported] - fission yeast			
Į		(Schizosaccharomyces pombe) emb CAB61771.1  (AL133225) probable pseudouridylate synthase			
3980	11359267		1.7		
3980	11339207	[Schizosaccharomyces pomoe]	1.7		
		gi 13652498 ref XP_007053.3  tubby like protein 3			
3982	13652498	[Homo sapiens] ref[XP_045148.1  8937 [Homo sapiens]	5.1		
		Transport of the second section of the second sections			
		gi 9978891 sp P57059 SN1L HUMAN PROBABLE			
		SERINE/THREONINE PROTEIN KINASE SNF1LK			
		dbj BAA95536.1  (AP001751) gene similar to rat			
3983	9978891	protein kinase (KID2) [Homo sapiens]	1.6		
		gi 7448436 pir  E71674 hypothetical protein RP682 -			
		Rickettsia prowazekii emb CAA15119.1  (AJ235272)			
3986	7448436	unknown [Rickettsia prowazekii]	2.2		
	 	gi 13815874 gb AAK42697.1  (AE006854) Hypothetical			
3987	13815874	protein [Sulfolobus solfataricus]	3.8		

are re		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
		gi 7485330 pir  T04802 hypothetical protein	
		F10M23.130 - Arabidopsis thaliana emb CAB36525.1	
		(AL035440) putative APG protein [Arabidopsis	
		thaliana] emb CAB79534.1  (AL161565) putative APG	
3989	7485330	protein [Arabidopsis thaliana]	4.5
		gi 7295073 gb AAF50399.1  (AE003555) Talin gene	
		product [Drosophila melanogaster] gb AAG22814.1	
3991	7295073		3.8
3371	1293013	gi 14456133 emb CAC41650.1  (AJ315577) putative	3.0
3992	14456122	nitrate reductase [Ustilago maydis]	8
3992	14430133	gi 533179 gb AAA56944.1  (L33090) pol protein	
3993	533179	[Human immunodeficiency virus type 2]	7.4
3993	333117	[Finnan manusodenestatey virus type 2]	7.4
		gi 14250408 gb AAH08638.1 AAH08638 (BC008638)	
3996	14250408	Similar to aspartyl-tRNA synthetase [Mus musculus]	4.1
		gi 14775307 ref XP_042250.1  similar to nuclear pore	
4000	14775307	complex interacting protein (H. sapiens) [Homo sapiens]	3.5
		gi 7507695 pir  T24847 hypothetical protein T11G6.7 -	
		Caenorhabditis elegans emb CAA93415.1  (Z69384)	
4002	7507695	predicted using Genefinder [Caenorhabditis elegans]	8.3
	1007070	promotes assigned (Sustained and Sustained	0.5
		gi 1173359 sp P45287 SAPC_HAEIN PEPTIDE	
		TRANSPORT SYSTEM PERMEASE PROTEIN	
		SAPC pir  C64134 sapC protein homolog -	
		Haemophilus influenzae (strain Rd KW20)	
		gb AAC23287.1  (U32837) peptide ABC transporter,	
4008	1173359	permease protein (sapC) [Haemophilus influenzae Rd]	0.83
		"12205520  Thip 000216 1  DIVER! DAIA	
		gi 13385538 ref NP_080316.1  RIKEN cDNA	
4009	12205520	2810036K01 gene [Mus musculus] dbj BAB28520.1	15.41
4009	13363338	(AK012865) putative [Mus musculus] gil13639013 refIXP 012007.2  dual specificity	1E-41
4011	13639013	phosphatase 8 [Homo sapiens]	9.1
7011	13039013	Procedumence of Liguide Subreness	9.1
		gi 1079359 pir  JC2394 phospholipase A2 inhibitor 25K	
		chain - monocled cobra gb AAB32583,1  phospholipase	
		A2 25 kda subunit, PLA2 25 kda subunit=urokinase-	
		type plasminogen activator receptor homolog [Naja naja	
4015	1079359	kaouthia=Thailand cobras, blood, Peptide, 185 aa]	5

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
4016	284079	gi 284079 pir  E41925 hypothetical protein 3 - human gb AAA58464.1  (M69297) ORF 3 [Homo sapiens]	0.005
4017	7293601	gi 7293601 gb AAF48973.1  (AE003512) CG14217 gene product [Drosophila melanogaster]	2.9
4020	11352313	gi 11352313 pir  G83376 probable trehalose synthase PA2152 [imported] - Pscudomonas acrugimosa (strain PAO1) gb AAG05540.1 AE004642_7 (AE004642) probable trehalose synthase [Pscudomonas acrugimosa]	2.9
4023	11386701	gi 11386701 sp Q9V773 C6AK_DROME PROBABLE CYTOCHROME P450 6A20 (CYPVIA20)	9
4030	7332064	gi[7332064 gb AAF60751.1  (AC006801) contains similarity to Loligo pealci microtubule-associated protein H1 (PIR:S28831) [Caenorhabditis elegans]	1.7
4034	13242494	gi 13242494 ref[NP_077507.1  EsV-1-22 [Ectocarpus siliculosus virus]	1.6
4035	7522108	gi 7522108 pir  T29097 pro-pol-dUTPase polyprotein - murine endogenous retrovirus ERV-L (fragment) emb[CAA73251.1] (Y12713) protease; reverse transcriptase; RNascH; integrase; dUTPase; Pro-Pol- dUTPase polyprotein [Mus musculus]	0.025
4042	12045115	gi[12045115]ref[NP_072926.1] lipoprotein, putative [Mycoplasma genitalium] sp]P47502[Y260_MYCGE HYPOTHETICAL LIPOPROTEIN MG260 PRECURSOR pirl[G64228 hypothetical protein homolog MG260 - Mycoplasma genitalium gb]AAC71481.1] (U39705) lipoprotein, putative [Mycoplasma genitalium]	3.2
4044		gi]14249544 ref[NP_116223.1  hypothetical protein FLJ14751 [Homo sapiens] dbj[BAB55272.1] (AK027657) unnamed protein product [Homo sapiens]	0.27
4046	6474685	gi 6474685 dbj BAA87286.1  (AB027982) Protein kinase C-like 1 [Schizosaccharomyces pombe]	1.9
4049	1514669	gi 1514669 emb CAA87082.1  (Z46958) adenyl cyclase [Xenopus lacvis]	0.56
4050	1514669	gi 1514669 emb CAA87082.1  (Z46958) adenyl cyclase [Xenopus laevis]	0.85

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
		33503411011	1 VALUE
		gi 2499547 sp P77153 WZB ECOLI PROBABLE	
		LOW MOLECULAR WEIGHT PROTEIN-	
		TYROSINE-PHOSPHATASE WZB pir D64972	
		probable protein-tyrosine-phosphatase (EC 3,1,3,48)	
		wzb, low molecular weight - Escherichia coli	
		gb AAC77834.1  (U38473) putative acid phosphatase	
		[Escherichia coli] gb AAC75122.1  (AE000296)	
		probable protein-tyrosine-phosphatase [Escherichia coli	
		K12] gb AAG57121.1 AE005432_2 (AE005432)	
		probable protein-tyrosine-phosphatase [Escherichia coli	
		O157:H7 EDL933] dbj BAB36289.1  (AP002560)	
		probable protein-tyrosine-phosphatase [Escherichia coli	
4051	2499547	O157:H7]	2.8
( )		gi 14916565 sp Q9XHG2 FLS_MALDO FLAVONOL	
4050	1401 0505	SYNTHASE (FLS) gb AAD26261.1 AF119095_1	
4052	14916565	(AF119095) flavonol synthase [Malus x domestica]	3.8
4050	250005	gi 3769667 gb AAC64604.1  (AF093797) unknown	
4053	3769667	[Norwalk virus]	2.8
4060	14225002	gi 14335082 gb AAK59820.1  (AY037220) AT4g24060/T19F6 50 [Arabidopsis thaliana]	1.3
4060	14333082	gi 12838540 dbj BAB24237,1  (AK005786) putative	1.3
4061	12929540	[Mus musculus]	4.3
4001	12030340	gi 7549797 ref NP_035731.1  T lymphoma oncogene	4,3
		[Mus musculus] sp[P17408 TLM MOUSE TLM	
		PROTEIN (TLM ONCOGENE) pir  S10151	
		transforming protein tlm - mouse (strain balb/c)	
		emb CAA36859.1  (X52634) tlm protein [Mus	
4066	7549797	musculus]	0.58
		gi 11357726 pir  T51437 hypothetical protein F2G14 40	
		- Arabidopsis thaliana emb CAC01811.1  (AL391146)	
		putative protein [Arabidopsis thaliana]	
1		gb AAK49610.1 AF372894 1 (AF372894)	
		AT5g14920/F2G14_40 [Arabidopsis thaliana]	
		gb AAK74054.1  (AY045696) AT5g14920/F2G14_40	
4067	11357726	[Arabidopsis thaliana]	1.8
		gi 8218109 cmb CAB92762.1  (AL121883)	
l		dJ545K15.1.3 (novel protein similar to KIAA0512	
		(contains translation of cDNAs Em:AK000818 and	
4 <b>0</b> 70	8218109	Em:L20773) (isoform 3)) [Homo sapiens]	5.5

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins
SEQ ID	ACCESS	Toarest Polghoot (Diasta vs. 140h-Redundam	riouns)
NO	N	DESCRIPTION	P VALUE
		gi 13096854 gb AAH03224.1 AAH03224 (BC003224)	
		Similar to chromosome condensation 1-like [Mus	
4072	13096854	musculus]	9.1
		gi 103076 pir  B21124 Bkm-like sex-determining region	
	i	hypothetical protein CS314 - fruit fly (Drosophila	
4073	103076	melanogaster) (fragment)	0.3
		gi 14722732 ref XP_017441.2  hypothetical protein	
4074	14722732	FLJ21477 [Homo sapiens]	6.6
		gi 3378554 emb CAA76192.1  (Y16404) multidrug	
4076	3378554	resistance protein [Candida albicans]	0.33
	İ	gi 6983946 gb AAF34740.1 AF200327_1 (AF200327)	
4081	6983946	unknown [Plasmodium chabaudi]	8.5
		gi 5524754 emb CAB50786.1  (AJ011801) Rx protein	
4082	5524754		2.6
	!	gi 7959271 dbj BAA96029.1  (AB040938) KIAA1505	
4086	7959271	protein [Homo sapiens]	3E-58
		gi 130687 sp P27536 POST_XENLA POSTERIOR	
		PROTEIN pir∥A43784 Xpo protein - African clawed	
4087	130687	frog emb CAA41397.1  (X58487) Xpo [Xenopus laevis]	1.2
		gi 2406633 gb AAB70469.1  (AF005630) adenylyl	
4092	2406633	cyclase isoform DAC9 [Drosophila melanogaster]	7.3
		gi 13812388 ref NP_113506.1  cell division cycle 2	
	1	homolog [Guillardia theta] emb CAC27075.1	
4005		(AJ010592) cell division cycle 2 homolog [Guillardia	
4095	13812388	thetaj	9.7
		113 450 000 01 1 1 1 COMO 00 01 1 1 1 000 000 00 00 00 00 00 00 0	
4096		gi 14530932 gb AAG37978.2  (AC087081) Hypothetical	
4090	14330932	protein Y82E9BL.13 [Caenorhabditis elegans]	3.1
4097	14760316	gi 14760316 reffXP_045601.1  7324 [Homo sapiens]	0.83
4021	14700310	gri 147003 Tolten AF_043001.1  7324 [Homo sapiens]	0.83
		gi 7305361 ref[NP_038652.1  otogelin [Mus musculus]	
		pir  T42214 otogelin - mouse gb AAB96561.1  (U96411)	
4098	7305361	otogelin; MLEMP [Mus musculus]	1E-10
4070	_	gi 8953656 emb CAB96704.1  (AL360354) vir15, rpg 1.	12-10
		putative transmembrane protein, similar to rpg2, vir9	
		MW:34499 (294 aa), fasta scores: opt: 398, E(): 0.023,	
		26.0% identity in 289 aa overlap, and to vir2:	
		MW:35019 (292 aa), fasta scores: opt: 380, E(): 0,037,	
4103	8953656	28.0% identity in>	3.6
		gi 3659615 gb AAC61684.1  (AF052517) diguanylate	
4108	3659615	cyclase [Gluconacetobacter xylinus]	6.5
		-, ( // / / / / / / / / / / /	0.0

400 TD		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
		gi 11465871 ref[NP_066420.1  NADH dehydrogenase	
		subunit 2 [Ochromonas danica]	
4110	11465051	gb AAG18386.1 AF287134_11 (AF287134) NADH	
4110	11465871	dehydrogenase subunit 2 [Ochromonas danica]	9.3
		gi 13786586 ref NP_112718.1  ORF55 [Bacteriophage TP901-1] gb AAK38072.1  (AF304433) ORF55	
4112	12796596	[Bacteriophage TP901-1]	1.4
4112	13780380	gi 14743649 rcf XP 036116.1  KIAA0594 protein	1.4
4115	14743649	[Homo sapiens]	2E-46
1115	11713013	gi 7491991 pir T41608 hypothetical protein	215-40
		SPCC790.03 - fission yeast (Schizosaccharomyces	
		pombe) emb CAA21293.1  (AL031855) hypothetical	
4121	7491991	protein [Schizosaccharomyces pombe]	2.9
		[	
		gi 6647413 sp Q9ZE52 ALR RICPR ALANINE	
		RACEMASE pir F71718 alanine racemase (alr) RP095	
		- Rickettsia prowazekii emb CAA14565.1  (AJ235270)	
4122	6647413	ALANINE RACEMASE (alr) [Rickettsia prowazekii]	9.1
		gi 7293814 gb AAF49181.1  (AE003517) nes gene	
4125	7293814	product [Drosophila melanogaster]	7.7
		gi 7505316 pir  T23351 hypothetical protein K05D4.2 -	
		Caenorhabditis elegans emb CAB07252.1  (Z92804)	
		contains similarity to Pfam domain: PF01461 (7TM	
		chemoreceptor), Score=118.9, E-value=3.1e-32, N=1	
4128	7505316		9.3
		gi 4335719 gb AAD17397.1  (AC006248) putative	
4139	4225510	C3HC4-type RING zinc finger protein [Arabidopsis	
4139	4335719	thaliana]	5.1
		gi 9625775 ref NP_040024.1  Conserved herpesvirus	
		spliced gene [human herpesvirus 5]	
		sp P16732 VTER_HCMVA PROBABLE DNA	
		PACKAGING PROTEIN emb CAA35363.1  (X17403) Conserved herpesvirus spliced gene [human herpesvirus	
4140	9625775	5]	4.3
	- 025,75		7.5
		gi 7510388 pir T27298 hypothetical protein Y68A4A,7	
		Caenorhabditis elegans emb CAA16418.1  (AL021503)	
		predicted using Genefinder~contains similarity to Pfam	
		domain: PF01604 (7TM chemoreceptor), Score=-48.3.	
4144	7510388	E-value=7.1e-07, N=1 [Caenorhabditis elegans]	3.4

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO NO	N	DESCRIPTION	P VALUE
		gi 7243081 dbj BAA92588.1  (AB037771) KIAA1350	
4145	7243081	protein [Homo sapiens]	e-112
		gi 13812221 ref NP_113352.1  hypothetical protein	
		[Guillardia theta] gb AAK39908.1 AF165818 116	
4149	13812221	(AF165818) hypothetical protein [Guillardia theta]	5.1
		gi 7706557 ref NP_057604.1  hepatocellular carcinoma-	
1	ł	associated antigen 59; hypothetical protein [Homo	
		sapiens] ref[XP_017552.1  hypothetical protein [Homo	
		sapiens] ref[XP_052691.1] hypothetical protein [Homo	
		sapiens] gb[AAF37561.1] (AF218421) hepatocellular	
		carcinoma-associated antigen 59 [Homo sapiens]	
		gb AAH07664.1 AAH07664 (BC007664) hepatocellular	
4150	7706557	carcinoma-associated antigen 59 [Homo sapiens]	3E-81
1150	7700337	gi 984305 gb AAA75468,1  (U25975) hPAK65 [Homo	515-01
4151	984305	sapiens]	1E-11
1101	301303	gi 14729667 ref XP 029101.1  KIAA0947 protein	115-11
4153	14729667	[Homo sapiens]	0.47
4133	14723007	gi 3355648 emb CAA08785.1  (AJ009688) tungsten	0.47
		formylmethanofuran dehydrogenase subunit fwdB	
4154	3355648	[Methanothermobacter wolfeii]	0.54
7134	3333040	[ivenianonermobacter wonen]	0.34
		-1125121021-114 AC54704 114 E005215 1 (4 E005215)	
		gi 12513192 gb AAG54704.1 AE005215_1 (AE005215) putative transport protein [Escherichia coli O157:H7	
		EDL933] dbj BAB33831.1  (AP002551) putative	
4156	12512102	transport protein [Escherichia coli O157:H7]	4,1
4130	12313192	gi 7472450 pir  D75370 hypothetical protein -	4,1
		Deinococcus radiodurans (strain R1) gb AAF11218.1 AE002008 13 (AE002008)	
4159	7472450	hypothetical protein [Deinococcus radiodurans]	7.9
4139	7472430		7.9
4160	12076525	gi 13876525 gb AAK43501.1 AC020666_11	
4100	136/0323	(AC020666) hypothetical protein [Oryza sativa]	4.6
4162	((6500	gi 6665675 gb AAF22966.1 AF175308_1 (AF175308)	OF 11
4162	6665675	acetyl-CoA carboxylase [Sus scrofa]	2E-11
		gi 7509105 pir  T31728 probable cysteine proteinase	
		(EC 3.4.22) W07B8.4 - Cacnorhabditis elegans	
4162	## OO 1 C 2	gb AAB65345.1  (AF016426) Hypothetical protein	
4163	7509105	W07B8.4 [Caenorhabditis elegans]	1.4
i l			
1	ĺ	gi 7510169 pir  T31555 hypothetical protein Y53H1C.2 -	
	==1016	Caenorhabditis elegans emb CAB55064.1  (AL117201)	_
4164	7510169	predicted using Genefinder [Caenorhabditis elegans]	7

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	t Proteins)
SEQ ID	ACCESS		T
NO	N	DESCRIPTION	P VALUE
		gi 6683126 dbj BAA20797.2  (AB002337) KIAA0339	
4166	6683126		9.8
1		gi 7485930 pir  T10623 hypothetical protein	
		F21C20.160 - Arabidopsis thaliana emb CAB45847.1	
		(AL080254) putative protein [Arabidopsis thaliana]	
		emb CAB79081.1  (AL161553) putative protein	
4167	7485930	[Arabidopsis thaliana]	5.9
		gi 11348277 pir  C83019 conserved hypothetical protein	
		PA5019 [imported] - Pseudomonas aeruginosa (strain	
İ		PAO1) gb AAG08404.1 AE004914_5 (AE004914)	
		conserved hypothetical protein [Pseudomonas	
4171	11348277	aeruginosa]	3.5
		gi 1150480 emb CAA91110.1  (Z54312) hypothetical	
4172	1150480	protein [Lactobacillus sakei]	3.2
		gi 14776432 ref XP_050788.1  similar to KIAA0133	
4173	14776432	gene product (H. sapiens) [Homo sapiens]	0.3
		gi 13569915 ref NP_112205.1  amnionless protein	
	l	[Homo sapiens] gb AAK28532.1 AF328788_1	
4175	13569915	(AF328788) amnionless [Homo sapiens]	8.9
	ł		
		gi 6978481 ref NP_036905.1  a-kinase anchoring protein	
		[Rattus norvegicus] sp Q62924 AK11_RAT A KINASE	
		ANCHOR PROTEIN 11 (PROTEIN KINASE A	
		ANCHORING PROTEIN 11) (PRKA11) (A KINASE	
		ANCHOR PROTEIN 220 KDA) (AKAP 220)	
]		pir  T42732 A-kinase anchoring protein AKAP 220 - rat	
		gb AAB06559.1  (U48288) AKAP 220 [Rattus	
4180	6978481	norvegicus]	3.1
		gi 14738118 rcf XP_048649.1  Apobec-1	
		complementation factor; APOBEC-1 stimulating protein	
1		[Homo sapiens] emb CAB94754.1  (AJ272078)	
4187	14738118	APOBEC-1 stimulating protein [Homo sapiens]	1E-37
4100		gi 13183342 gb AAK15157.1 AF284034_1 (AF284034)	
4189	13183342	vitellogenin B [Melanogrammus aeglefinus]	9.4

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		Table 2D Market Control	
CEO ID		Table 3B Nearest Neighbor (BlastX vs. Non-Redundam	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
		gi 13123985 sp Q9N293 B3G5 GORGO BETA-1.3-	
		GALACTOSYLTRANSFERASE 5 (BETA-1,3-	
		GALTASE 5) (BETA3GAL-T5) (B3GAL-T5) (UDP-	
		GALACTOSE:BETA-N-ACETYLGLUCOSAMINE	
		BETA-1,3-GALACTOSYLTRANSFERASE 5) (UDP-	
		GAL:BETA-GLCNAC BETA-1,3-	
		GALACTOSYLTRANSFERASE 5) (BETA-3-GX-T5)	
		dbj BAA94501.1  (AB041416) UDP-Gal:GlcNAc	
4192	13123985	beta1,3-galactosyltransferase 5 [Homo sapiens]	4
4104	7205760	gi 7295768 gb AAF51070.1  (AE003579) CG3410 gene	
4194	7295768	product [Drosophila melanogaster] gi 7290544 gb AAF45996,1  (AE003432) CG6903 gene	1.5
4195	7290544	gi /290544 gb AAF45996.1  (AE003432) CG6903 gene product [Drosophila melanogaster]	
4193	7290344	gi 9966502 gb AAG10295.1 AF275272 1 (AF275272)	6
4200	9966502	unknown [Schizophyllum commune]	0.83
-1200	3300302	diadiewa [Bellizophynam commune]	0,63
		gi 6010435 gb AAF01135.1  (AF087699) erythrocyte	
4213		membrane protein 3 [Plasmodium falciparum]	2.2
		gi 14775884 ref XP 043964.1  similar to KIAA0220	
4216	14775884	protein (H. sapiens) [Homo sapiens]	5E-12
		gi 7511578 pir T19209 probable protein kinase	
		E02H4.3 - Caenorhabditis elegans emb CAA94122.1	
		(Z70205) Similarity to Drosophila Doa kinase (PIR	
		Acc. No. S44077), contains similarity to Pfam domain:	
		PF00069 (Eukaryotic protein kinase domain),	
		Score=184.6, E-value=5.2e-52, N=1~cDNA EST	
		yk5d6.3 comes from this gene~cDNA EST yk5d6.5	
		comes from this> emb CAA91979.1  (Z68003)	
		Similarity to Drosophila Doa kinase (PIR Acc. No.	
		S44077), contains similarity to Pfam domain: PF00069	
		(Eukaryotic protein kinase domain), Score=184.6, E- value=5.2e-52, N=1~cDNA EST yk5d6.3 comes from	
4218		this gene~cDNA EST yk5d6.5 comes from this>	0.069
7210		gi 7471938 pir  G75366 glucose-1-phosphate	0,009
		adenylyltransferase - Deinococcus radiodurans (strain	
		R1) gb AAF11244.1 AE002010 7 (AE002010) glucose-	,
		1-phosphate adenylyltransferase [Deinococcus	
4220	7471938	radioduransl	2.6

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	· Dectains)
SEQ ID	ACCESS	Table 55 Postest Neighbor (Diasta vs. Non-Redundam	riotems)
NO	N	DESCRIPTION	P VALUE
l		gi 4506051 ref NP 000937.1  primase, polypeptide 1	
		(49kD); primase polypeptide 1 (49kD) [Homo sapiens]	
		ref[XP_048930.1] primase, polypeptide 1 (49kD) [Homo	
		sapiens] ref[XP_006706.3] primase, polypeptide 1	
		(49kD) [Homo sapiens] sp[P49642[PRI1_HUMAN	
		DNA PRIMASE SMALL SUBUNIT (DNA PRIMASE	
ì	1	49 KDA SUBUNIT) (P49) pir  S45630 DNA primase	
		chain p48 - human emb CAA52377.1  (X74330) DNA	
		primase (subunit p48) [Homo sapiens]	
		gb AAH05266.1 AAH05266 (BC005266) primase,	
4224	4506051	polypeptide 1 (49kD) [Homo sapiens]	0.014
	İ	gi 13812048 ref NP 113182.1  putative protein kinase	
		[Guillardia theta] gb AAK39750.1 AF083031 107	
4228	13812048		1.8
	15012010	gi 7465475 pir  S70173 autoinducer synthesis protein -	1.0
4229	7465475	Pseudomonas aeruginosa	1.3
		gi 11288518 pir  T49586 related to nif-specific	
İ		regulatory protein [imported] - Neurospora crassa	
		emb CAB91387.1  (AL355930) related to nif-specific	
4236	11288518	regulatory protein [Neurospora crassa]	6.8
		gi 2072951 gb AAC51263.1  (U93564) putative p150	
4237	2072951	[Homo sapiens]	3.1
		gi 1903416 gb AAC53095.1  (U76112) translation	
4244	1903416	repressor NAT1 [Mus musculus]	2E-15
4248	14740171	gi 14749171 ref XP_038309.1  hypothetical protein XP 038309 [Homo sapiens]	4.2
4240	14/491/1		4.3
ŀ	1	gi 7432367 pir  T13881 NADH dehydrogenase (ubiquinone) (EC 1.6.5.3) chain 4 - Chlorogonium	
		elongatum mitochondrion emblCAA73990.11 (Y13644)	
ĺ		NADH dehydrogenase subunit 4 [Chlorogonium	
4251	7432367	elongatum]	1.2
		gi 13621755 gb AAK33536,1  (AE006511) putative	
]	1	nucleotide sugar dehydrogenase [Streptococcus	
4255	13621755	pyogenes M1 GAS]	2.7
		gi 9880 emb CAA36427.1  (X52177) RNA polymerase	
4259	9880	beta subunit (697 AA) [Plasmodium falciparum]	1.1
		gi 10314010 ref[NP_066241.1  replicase polyprotein	
		[acute bee paralysis virus]	
4260	10214010	gb AAG13118.1 AF150629_1 (AF150629) replicase	
4260	10514010	polyprotein [acute bee paralysis virus]	7.5

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r		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		2 2000110)
NO	N	DESCRIPTION	P VALUE
		gi 1170902 sp P43245 MDR1 RAT MULTIDRUG	
		RESISTANCE PROTEIN 1 (P-GLYCOPROTEIN 1)	
4261	1170902	pir  JH0502 p-glycoprotein - rat	5,2
		gi 10180785 gb AAG14273.1 AF243438 88	
	1	(AF243438) RS1 immediate-early gene transactivator-	
		like protein [Gallid herpesvirus 2]	
		gb AAG14284.1 AF243438 99 (AF243438) RS1	
		immediate-early gene transactivator ICP4-like protein	
4265	10180785	[Gallid herpesvirus 2]	1.3
		gi 1353257 gb AAB06234.1  (U26665) dimethyl	
		sulphoxide reductase subunit B [Haemophilus	
4267	1353257	influenzae]	4.8
		gi 4240315 dbj BAA74936.1  (AB020720) KIAA0913	
4268	4240315	protein [Homo sapiens]	3.3
		gi 4493991 cmb CAB39050.1  (AL034559) hypothetical	
4269	4493991	protein, PFC1060c [Plasmodium falciparum]	6.3
		gi 4557503 ref NP_001072.1  cubilin precursor; cubilin;	
		intrinsic factor-cobalamin receptor; intrinsic factor B12-	
		receptor [Homo sapiens] pir  T09456 intrinsic factor-	
		B12 receptor Cubilin precursor - human	
		gb AAC82612.1  (AF034611) intrinsic factor-B12	
4272	4557503	receptor precursor; cubilin [Homo sapiens]	5E-10
		gi 7504212 pir  T22670 hypothetical protein F54F11.1 -	
		Caenorhabditis elegans emb CAB05738,1  (Z83229)	
		Weak similarity with glycerol phospholipid-cholesterol	
		acyltransferase~cDNA EST yk479h12.3 comes from	
		this gene~cDNA EST yk479h12.5 comes from this gene	
4277	7504212	[Caenorhabditis elegans]	8
		gi 421548 pir  B47013 butanol dehydrogenase (EC 1.1.1.	
4279	421548	) II - Clostridium acetobutylicum	1.7
	l	gi 7497553 pir  T19999 hypothetical protein C47D12.2 -	
		Caenorhabditis elegans cmb CAA93767.1  (Z69902)	
		predicted using Genefinder~cDNA EST yk14e2.3 comes	-
		from this gene~cDNA EST yk88e3.3 comes from this	
	ĺ	gene~cDNA EST yk132h1.3 comes from this	
		gene~cDNA EST yk88e3.5 comes from this	
		gene~cDNA EST yk132h1.5 comes from this	
4284	7497553	gene~cDNA EST yk422c3.3 >	6.7

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	
NO	- N	DESCRIPTION	P VALUE
i		"Southern the Public of the Company	
]		gi 5042237 emb CAB44655.1  (Y18605) hypothetical	
İ		protein RvD1-Rv2024c' [Mycobacterium bovis BCG]	
		gb AAK46361.1  (AE007059) helicase, putative/conserved hypothetical protein [Mycobacterium]	
4292	5042237	tuberculosis CDC15511	
4292	3042237	tuberculosis CDC1551]	6,3
1		gi 14746217 ref XP 011563.2  hypothetical protein	
		FLJ21634 [Homo sapiens] dbj[BAB15338.1]	
4298	14746217	(AK026056) unnamed protein product [Homo sapiens]	0.000000002
4230	14740217	gi 6005978 ref NP 009098.1  zinc finger protein 258	0.00000002
		[Homo sapiens] gb[AAD15797.1] (AF055470) ZNF258	
4299	6005978		9E-26
4233	0003976		9E-20
		gi 9845291 ref NP_063945.1  phosphatidylinositol polyphosphate 5-phosphatase type IV [Homo sapiens]	
		gb AAF81404.1 AF187891 1 (AF187891)	
		phosphatidylinositol polyphosphate 5-phosphatase type	
4303	09/5201	IV [Homo sapiens]	7.7
4303	9043291	gi 13620879 dbi BAB40992.1  (AB049938) alpha1,3-	
4305	13620879	fucosyltransferase [Rattus norvegicus]	2.7
4303	13020073	gi 6424813 gb AAF08148.1  (AF130192) NADH	2.1
4308	6424813	dehydrogenase subunit F [Valeriana fauriei]	9.6
		gi 5834894 ref NP 006964.1 ND5 10021 NADH	
		dehydrogenase subunit 5 [Caenorhabditis elegans]	
		sp P24896 NU5M CAEEL NADH-UBIQUINONE	
		OXIDOREDUCTASE CHAIN 5 pir    \$26037 NADH	
		dehydrogenase (ubiquinone) (EC 1.6.5.3) chain 5 -	
		Caenorhabditis elegans mitochondrion	
		emb CAA38162.1  (X54252) ND5 protein (AA 1 - 527)	
4311	5834894	[Caenorhabditis elegans]	1.6
		gi 11135954 sp Q9TC94 YM16_NEPOL	
		HYPOTHETICAL PROTEIN YMF16	
		gb AAF03203.1 AF110138_35 (AF110138) homolog of	
		E. coli MttB, a protein involved in folded protein	
		translocation and targeting across bacterial membranes	
4314	11135954	[Nephroselmis olivacea]	2.2
		- 1140909211 - 11G1-G12500 11 (AT 4455C D - 1	
4316	14000021	gi 14089831 emb CAC13590.1  (AL445564) unknown; predicted coding region [Mycoplasma pulmonis]	6.5
4510	14069831	gi 10728595 gb AAF52302,2  (AE003611) CG9011	6.5
4317	10728595	ge 10/28595 gb AAF52302.2  (AE003611) CG9011 gene product [Drosophila melanogaster]	4E-19
7311	10/20093	gene product (Diosophila meianogaster)	46-19

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
	}	701074701 11/74704040 11/44007700	
4318	9187479	gi 9187479 emb CAB96962.1  (AJ007508) magnesium- chelatase subunit chll [Gnetum gnemon]	2.4
4318	918/4/9	chelatase subunit chii [Gnetum gnemon]	2.4
		gi 7469635 pir  \$76563 hypothetical protein -	
		Synechocystis sp. (strain PCC 6803) dbi BAA10409.1	
		(D64002) ORF ID:sll0188~unknown protein	
4322	7469635	[Synechocystis sp. PCC 6803]	1.6
		gi 15021476 gb AAK77753.1 AF369029 84	
4324	15021476	(AF369029) ORF84 [white spot syndrome virus]	1.7
		gi 11466210 ref NP_066533.1  NADH dehydrogenase	
		subunit 2 [Naegleria gruberi]	
		gb AAG17811.1 AF288092_36 (AF288092) NADH	
4333	11466210	dehydrogenase subunit 2 [Naegleria gruberi]	0.46
		gi 127494 sp P25127 MT_ESOLU	
		METALLOTHIONEIN (MT) pir   S17175	
	İ	Metallothionein - Northern pike pir  S38334 metallothionein - northern pike pir  S31723	
		metallothionein - northern pike emb CAA42035,1	
	Į.	(X59392) metallothionein [Esox lucius]	
		emb CAA49636.1  (X70042) Metallothioein [Esox	
4336	127494	lucius]	1
		gi 6009729 dbj BAA85039.1  (AB026129) alpha-2-	
4341	6009729	macroglobulin-2 [Cyprinus carpio]	0.00002
•		gi 1170606 sp P43188 KADC_MAIZE ADENYLATE	
	ì	KINASE, CHLOROPLAST (ATP-AMP	
		TRANSPHOSPHORYLASE) pir  S45634 adenylate	
		kinase (EC 2.7.4.3), chloroplast - maize pdb IZAK A	
		Chain A, Adenylate Kinase From Maize In Complex	
		With The Inhibitor P1,P5-Bis(Adenosine-5'- )pentaphosphate (Ap5a) pdb 1ZAK B Chain B,	
		Adenylate Kinase From Maize In Complex With The	
		Inhibitor P1,P5-Bis(Adenosine-5'-)pentaphosphate	
4342	1170606	(Ap5a)	5E-13
		gi 543537 pir  JN0720 glucosyl transferase - Shigella	
	1	flexneri phage X gb AAA71895.1  (L05001) glucosyl	
4345	543537	transferase [bacteriophage SfX]	8.1
		gi 7297011 gb AAF52281.1  (AE003611) CG11149	
4347	7297011	gene product [Drosophila melanogaster]	0.93
1210		gi 11558496 cmb CAC17816.1  (AJ276292) sodium	
4348	L11558496	iodide symporter [Sus scrofa]	2.6

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 7293017 gb AAF48404.1  (AE003497) CG12398	
4351	7293017	gene product [Drosophila melanogaster]	9
		gi 7486021 pir  T06133 hypothetical protein	
	,	F23E12.200 - Arabidopsis thaliana emb CAA18745.1	
		(AL022604) putative protein [Arabidopsis thaliana]	
1		emb CAB80241.1  (AL161587) putative protein	
4352	7486021	[Arabidopsis thaliana]	0.085
		gi 10047323 dbj BAB13449.1  (AB046843) KIAA1623	
4353	10047323	protein [Homo sapiens]	2.7
		gi 262249 gb AAB24620.1  (S52010) orf1 5' of EpoR	
4359	262249	[Mus sp.]	0.67
		gi 14589828 gb AAK70656.1 AC024771 1	
		(AC024771) Hypothetical protein Y40B10A.1	
4361	14589828	[Caenorhabditis elegans]	0.45
		gi 543537 pir  JN0720 glucosyl transferase - Shigella	
		flexneri phage X gb AAA71895.1  (L05001) glucosyl	
4364	543537	transferase [bacteriophage SfX]	6.6
		gi 14587208 dbj BAB61142.1  (AP003199) hypothetical	
4366	14587208	protein [Oryza sativa]	5.1
		gi 13385738 ref NP_080509.1  RIKEN cDNA	
		4933434I20 gene [Mus musculus] dbj BAB30173.1	
4370	13385738	(AK016272) putative [Mus musculus]	0.046
		gi 7507823 pir  T24908 hypothetical protein T14D7.2 -	
		Caenorhabditis elegans emb CAB03365.1  (Z81123)	
		cDNA EST EMBL:M88978 comes from this	
		gene~cDNA EST yk419b8.3 comes from this	
		gene~cDNA EST yk285a8.3 comes from this	
		gene~cDNA EST yk398c1.3 comes from this	
		gene~cDNA EST yk350c8.3 comes from this	
4372	7507823	gene~cDNA EST yk350c8.5 comes from this gene~c>	0.33
		gi 7302573 gb AAF57655.1  (AE003798) CG15086	
4376	7302573	gene product [Drosophila melanogaster]	1.1
		gi 531764 emb CAA56961.1  (X81072) hypothetical	
4377	531764	protein in YTA7 5'region [Saccharomyces cerevisiae]	5.4

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundam	t Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 585100 sp Q07287 ZPB_PIG ZONA PELLUCIDA	
	l	SPERM-BINDING PROTEIN B PRECURSOR	
		(ZONA PELLUCIDA 3-ALPHA) (ZONA	
ļ		PELLUCIDA GLYCOPROTEIN ZP3-ALPHA)	]
		pir  S35712 sperm-binding glycoprotein ZP3-alpha	
		precursor - pig gb AAA50164.1  (L11000) zp3-alpha	
4378	585100	sperm-binding glycoprotein [Sus scrofa]	2.5
		gi 2148043 pir  S51527 S-receptor kinase (EC 2.7.1,-)	
		A14 precursor - rape gb AAA62232.1  (U00443) S-	
4382	2148043	receptor kinase [Brassica napus]	6
		gi 2257556 dbj BAA21448.1  (AB004539) probable	
		membrane protein YOL130w [Schizosaccharomyces	
4383	2257556	pombe]	7.9
		gi 8928528 sp Q44601 YTR1_BUCSC	
		HYPOTHETICAL 35.1 KDA PROTEIN IN TRPA	
		3'REGION gb AAA92793.1  (U09185) unknown	
4386	8928528	[Buchnera aphidicola]	9.3
		gi 12518050 gb AAG58518.1 AE005564_3 (AE005564)	
		maltodextrin phosphorylase [Escherichia coli O157:H7	
		EDL933] dbj BAB37682.1  (AP002565) maltodextrin	
4389	12518050	phosphorylase [Escherichia coli O157:H7]	4.8
		gi 13813161 gb AAK40397.1  (AE006644) Hypothetical	
4391	13813161	protein [Sulfolobus solfataricus]	0.64
		gi 13626090 sp P78704 ACR2_NEUCR	
		ACRIFLAVINE SENSITIVITY CONTROL	
4393	13626090	PROTEIN ACR-2	3
		gi 6680736 ref NP_031522.1  AT motif binding factor 1	
		[Mus musculus] dbj BAA05046.1  (D26046) AT motif-	
		binding factor [Mus musculus] prf  2207230A	
4394	6680736		1.1
		gi 7269957 emb CAB79774.1  (AL161577) cyclic	
		nucleotide and calmodulin-regulated ion channel-like	
4395	7269957	protein [Arabidopsis thaliana]	5
		gi 13812152 ref[NP_113279.1  hypothetical protein	
		[Guillardia theta] gb AAK39839.1 AF165818_47	
4396	13812152	(AF165818) hypothetical protein [Guillardia theta]	2

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
1		1112500741 185007411 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	
		gi 11350074 pir  G83141 hypothetical protein PA4035	
ì		[imported] - Pseudomonas aeruginosa (strain PAO1) gb AAG07422.1 AE004820_7 (AE004820) hypothetical	1
4398	11250074	protein [Pseudomonas acruginosa]	1.0
4376	11330074	protein [Fseudomonas acruginosa]	1.8
		gi 6601484 gb AAF18995,1 AF211856 1 (AF211856)	
		pulmonary surfactant protein A [Ovis aries]	
1		gb AAF31148.1 AF076633 1 (AF076633) pulmonary	
4399	6601484	surfactant-associated protein A [Ovis aries]	2.3
ì		gi 7480194 pir  T37054 hypothetical protein SCJ21.05 -	
		Streptomyces coelicolor emb CAB52351.1  (AL109747)	
4402	7480194	hypothetical SCJ21.05 [Streptomyces coelicolor A3(2)]	6.9
		gi 123346 sp P16393 HMDH STRPU 3-HYDROXY-3-	
		METHYLGLUTARYL-COENZYME A	
		REDUCTASE (HMG-COA REDUCTASE)	
		pir  A31898 hydroxymethylglutaryl-CoA reductase	
		(NADPH) (EC 1.1.1.34) - sea urchin	
		(Strongylocentrotus purpuratus) gb AAA30060.1	
1		(J04200) HMGCoA reductase (EC 1.1.1.34)	
4403	123346	[Strongylocentrotus purpuratus]	4.2
		gi 1870163 emb CAB05927.1  (Z83335) unknown	
4404	1870163	[Streptococcus pneumoniae]	8.3
		gi 5739524 ref NP_000544.1  Werner syndrome protein	
		[Homo sapiens] sp Q14191 WRN_HUMAN WERNER	
		SYNDROME HELICASE gb AAC41981.1  (L76937)	
[		Homo sapiens Werner syndrome gene, complete cds	
4400	£220524	gb AAC63361.1  (AF091214) WRN [Homo sapiens]	
4408	5739524		7.2
		gi 7494315 pir  C71607 hypothetical protein PFB0745w	
		malaria parasite (Plasmodium falciparum) gb[AAC71938.1] (AE001415) hypothetical protein	
4409	7494315		2.4
4409	1474313	gi 13477147 gb AAH05031.1 AAH05031 (BC005031)	2.4
		Unknown (protein for IMAGE:3532103) [Homo	
4414	13477147		9.1
4414	154//14/	gi 6650234 gb AAF21778.1  (AF072567) zinc finger	9.1
4415	6650234	protein 74 isoform III [Homo sapiens]	3.3
7715	5050254	protona / / isotorin in promo supions]	ــــــــــــــــــــــــــــــــــــــ

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NO   N   DESCRIPTION   P VALUE			Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
gij17498637 pir  T32474 hypothetical protein F08F1.3 - Caenorhabditis elegans gb]AAB71303.1  (AF026213)     4419	SEQ ID	ACCESS	DESCRIPTION	DAVALIE
Caenorhabditis elegans gb AAB71303.1  (AF026213)	- 110		DISCRIPTION	I VALUE
4419		1	gi 7498637 pir  T32474 hypothetical protein F08F1.3 -	
\$\frac{\text{gi}[6016842]\text{dbj}[BAA85182.1]}{\text{AB033168}\text{nuclear}}				
4420   6016842   protein ZAP [Mus musculus]   2.2	4419	7498637		9.9
gij125704 sp P08630 SRC2_DROME TYROSINE-PROTEIN KINASE SRC28C pir[ITVFFDS protein-tyrosine kinase (EC 2.7.1.112) src2 - fruit fly (Drosophila melanogaster) gb AAA28912.11 (M16599)				
PROTEIN KINASE SRC28C pir  TVFFDS protein-tyrosine kinase (BC 2.7.1.1) psr2 - fruit fly (Drosophila melanogaster) gb AAA5912.1  (M16599)	4420	6016842	protein ZAP [Mus musculus]	2.2
PROTEIN KINASE SRC28C pir  TVFFDS protein-tyrosine kinase (BC 2.7.1.1) psr2 - fruit fly (Drosophila melanogaster) gb AAA5912.1  (M16599)			The square process of the process of	
tyrosine kinase (EC 2.7.1.112) src2 - fruit fly (Drosophila melanogaster) gljAAA28912.11 (M16599) 4422 125704 Dsrc28C protein [Drosophila melanogaster] 4423 14746157 [Elomo sapiens] 4424 13235641 [Elomo sapiens] 4424 13235641 [Stephanotis floribunda] 4426 13235641 [Stephanotis floribunda] 4426 13235641 [Stephanotis floribunda] 4427 13235641 [Stephanotis floribunda] 4428 13235641 [Stephanotis floribunda] 4429 1428 7488979 [pit][T07612 cellulase (EC 3.2.1.4) Cel3, membrane-anchored - tomato gb]AAC49704.1] (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (Lycopersicon (U78526) endo-1,4-beta-glucanase (U78526) endo-1,4-beta-glucanase (U78526) endo-1,4-beta-glucanase (U78526) endo-1,4-beta-glucanase (U78526) endo-1,4-beta-glucanase (U78526) endo-1,4-beta-glucanase (U78526) endo-1,4-beta-glucanase (U78526) endo-1,4-beta-glucanase (U78526) endo-1,4-beta-glucanase (U78526) endo-1,4-beta-glucanase (U78526) endo-1,4-beta-glucanase				}
Crosophila melanogaster) gb[AAA28912.1] (M16599)   8.9				}
4422   125704   Darc28C protein [Drosophila melanogaster]   8,9				
14746157   Editor	4422	125704		8.9
gji 3235641 emb CAC33768.1  (AJ308570) S-adenosyl- L-methionine:salicylic acid carboxyl methyltransferase   4424   13235641 [Stephanotis floribunda]   6.3     gji 2805257 gb AAH02091.1 AAH02091 (BC002091)     4426   12805257   Unknown (protein for MGC:6342) [Mus musculus]   5.5     gji 7488979 pir [T07612 cellulase (BC 3.2.1.4) Cel3, membrane-anchored - tomato gb AAC49704.1] (U78526) endo-1,4-beta-glucanase [Lycopersicon   9.5     4428   7488979   esculentum]   9.5     4429   14741807   [Homo sapiens]   2E-16     4429   14741807   [Homo sapiens]   2E-16     25   26   26   27   27   27   27     26   27   27   27   27   27   27     27   27				
L-methionine:salicylic acid carboxyl methyltransferase   6.3	4423	14746157	[Homo sapiens]	8.6
12805257 gb AAH02091.1 AAH02091 (BC002091)			L-methionine:salicylic acid carboxyl methyltransferase	
12805257   Unknown (protein for MGC:6342) [Mus musculus]   5.5	4424	13235641	[Stephanotis floribunda]	6.3
12805257   Unknown (protein for MGC:6342) [Mus musculus]   5.5				
gij7488979 pirj[T07612 cellulase (EC 3.2.1.4) Cel3, membrane-anchored - tomato gbjAcC49704.1] (U78526) endo-1,4-beta-glucanase [Lycopersicon esculentum] 9,5   gij4741807 ref[XP_049622.1] KIAA1630 protein   9,5   gij6324137 ref[NP_014207.1] chitin synthase 1; Chs1p   Saccharomyces cerevisine] spjP08004 CHS1_YEAST   CHITIN SYNTHASE 1 (CHITIN-UDP ACETYL-GLUCOSAMINYL TRANSFERASE 1) pinjlA23944 chitin synthase (EC 2.4.1.16) CHS1 - yeast (Saccharomyces cerevisine) spjP080404 CHS1_YEAST   CHITIN-UDP ACETYL-GLUCOSAMINYL TRANSFERASE 1) pinjlA23944   Chitin synthase (EC 2.4.1.16) CHS1 - yeast (Saccharomyces cerevisine) gbjAAA34491.1] (M14045)				
membrane-anchored - tomato gb AAC49704.1    (U78526) endo-1,4-beta-glucanase [Lycopersicon   signatum   9,5     14741807 [reffXP_049622.1] KIAA1630 protein   14741807 [reffXP_049622.1] KIAA1630 protein   gi 6324137 [reffXP_014207.1] chitin synthase 1; Chs 1p   Saccharomyces cerevisine] spp(08004[CHS1_YEAST]   CHITIN SYNTHASE 1; (CHITIN-UDP ACETYL-   GLUCOSAMINYL TRANSFERASE 1) pir  A23944    chitin synthase (EC 2.4.1.16) CHS1 - yeast   Saccharomyces cerevisine] spl(AAA34491.1] (M14045)	4426	12805257		5.5
(178526) endo-1,4-beta-glucanase [Lycopersicon seculentum]   9,5				
4428   7488979   esculentum]   9.5				
gijl4741807 refjXP_049622.1  KIAA1630 protein  2E-16  gij6324137 reffNP_014207.1  chitin synthase 1; Chs1p [Saccharomyces cerevisiae] sppP08004 CHS1_YEAST CHITIN SYNTHASE 1 (CHITIN-UDP ACETYL- GLUCOSAMINYL TRANSFERASE 1) pirjlA23944 chitin synthase (EC 2.4.1.16) CHS1 - yeast (Saccharomyces cerevisiae) gbJAAA34491.1  (M14045)	4428	7488979		9.5
4429 14741807 [Homo sapiens] 2E-16  gij6324137[ref[NP_014207.1] chitin synthase 1; Chs1p [Saccharomyces cerevisiae] spjP08004[CHS1_YEAST CHITIN SYNTHASE 1 (CHITIN-UDP ACETYL- GLUCOSAMINYL TRANSFERASE 1) pirjlA23944 chitin synthase (EC 2.4.1.16) CHS1 - yeast (Saccharomyces cerevisiae) gbJAAA34491.1] (M14045)	7720	7400575		7.5
[Saccharomyces cerevisiae] sp[P08004 CHS1_YEAST CHITIN SYNTHASE 1 (CHITIN-UDP ACETYL- GLUCOSAMINYL TRANSFERASE 1) pirl A23944 chitin synthase (EC 2.4.1.16) CHS1 - yeast (Saccharomyces cerevisiae) gb AA33491.1  (M14045)	4429	14741807		2E-16
[Saccharomyces cerevisiae] sp[P08004 CHS1_YEAST CHITIN SYNTHASE 1 (CHITIN-UDP ACETYL- GLUCOSAMINYL TRANSFERASE 1) pirl A23944 chitin synthase (EC 2.4.1.16) CHS1 - yeast (Saccharomyces cerevisiae) gb AA33491.1  (M14045)				
[Saccharomyces cerevisiae] sp[P08004 CHS1_YEAST CHITIN SYNTHASE 1 (CHITIN-UDP ACETYL- GLUCOSAMINYL TRANSFERASE 1) pirl A23944 chitin synthase (EC 2.4.1.16) CHS1 - yeast (Saccharomyces cerevisiae) gb AA33491.1  (M14045)			gil6324137lrefINP 014207.11 chitin synthase 1: Chs1n	
CHITIN SYNTHASE 1 (CHITIN-UDP ACETYL- GLUCOSAMINYL TRANSFERASE 1) pirilA23944 chitin synthase (EC 2.4.1.16) CHS1 - yeast (Saccharomyces cerevisiae) gbJAAA34491.1  (M14045)	1			
chitin synthase (EC 2.4.1.16) CHS1 - yeast (Saccharomyces cerevisiae) gb[AAA34491.1] (M14045)				
(Saccharomyces cerevisiae) gb AAA34491.1  (M14045)			GLUCOSAMINYL TRANSFERASE 1) pir A23944	
			chitin synthase (EC 2.4.1.16) CHS1 - yeast	
chitin synthase [Saccharomyces cerevisiae]				
			chitin synthase [Saccharomyces cerevisiae]	
emb CAA96086.1  (Z71468) ORF YNL192w  4434   6324137   [Saccharomyces cerevisiae]   6.8	4424	6204127		
	4434	6324137		6.8
gi 6678958 ref NP_032667.1  microtubule associated				
testis specific serine/threonine protein kinase [Mus musculus] pir  A54602 microtubule-associated				
serine/threonine protein kinase MAST205 - mouse				
gb AAC04312.1  (U02313) protein kinase [Mus				
4436   6678958   musculus] 0.41	4436	6678958		0.41

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 7494262 pir  T18489 hypothetical protein C0820w -	
		malaria parasite (Plasmodium falciparum)	
		emb CAB11128.1  (Z98551) hypothetical protein,	
4445	7494262	PFC0820w [Plasmodium falciparum]	6.3
	4		
		gi 11245201 gb AAG33502.1 AF302058_1 (AF302058)	
4454	11245201	cytochrome oxidase I [Apocrypta sp. MSPAR2]	9.1
		gi 6752960 ref[NP 033743.1  a disintegrin and	
		metalloprotease domain 11; a disintegrin and	
		metalloprotease domain (ADAM) 11 [Mus musculus]	
		sp Q9R1V4 AD11 MOUSE ADAM 11 PRECURSOR	
		(A DISINTEGRIN AND METALLOPROTEINASE	
		DOMAIN 11) (METALLOPROTEINASE-LIKE.	
		DISINTEGRIN-LIKE, AND CYSTEINE-RICH	
		PROTEIN) (MDC) dbj BAA83384.1  (AB009676)	
4457	6752960		3.8
	0,02,00	gi 103076 pir  B21124 Bkm-like sex-determining region	3.0
		hypothetical protein CS314 - fruit fly (Drosophila	
4462	103076	melanogaster) (fragment)	0.14
	100070	gi 14742023 ref XP 039778,1  HSPC047 protein [Homo	0.14
4463	14742023		1E-65
1105	11712020	gi 13816057 gb AAK42842.1  (AE006867) Agmatinase	112-05
		(agmatine ureohydrolase) (speB-2) [Sulfolobus	
4464	13816057	solfataricus]	6.5
	15510057	gi 295359 gb AAA21303.1  (L14824) surface antigen	0.5
4466	295359	[Trypanosoma cruzi]	4.2
		gi 2429459 gb AAB70995.1  (AF025461) contains	
		similarity to Canis familiaris (dog) 180k ribosome	
4471	2429459	receptor (NID:g984113) [Caenorhabditis elegans]	2.2
		gi 7296176 gb AAF51469.1  (AE003588) CG2839 gene	
4474	7296176		8.9
		gi 14736857 ref XP 005322.2  KIAA0967 protein	
4475	14736857	[Homo sapiens]	0.33
		gi 13959004 gb AAK51055.1 AF361075 2 (AF361075)	
4480	13959004	UL24 [Canine herpesvirus]	0.35

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	(See Long Long Long Long Long Long Long Long	
NO	N	DESCRIPTION	P VALUE
		gi 7511539 pir  T18770 probable calcium channel	
		protein - Caenorhabditis elegans emb CAA90091.1	
		(Z49907) similar to calcium channel alpha-2	
		subunit~cDNA EST yk134g4.5 comes from this	
		gene~cDNA EST yk49c7.3 comes from this	
		gene~cDNA EST yk134g4.3 comes from this	
		gene~cDNA EST yk49c7.5 comes from this	
		gene~cDNA EST yk349a4.3 comes from this gene~cD>	
		emb CAA90141.1  (Z49912) similar to calcium channel alpha-2 subunit~cDNA EST yk134g4.5 comes from this	
		gene~cDNA EST yk49c7,3 comes from this	
		gene~cDNA EST yk49c7,3 comes from this gene~cDNA EST yk134g4.3 comes from this	
		gene~cDNA EST yk194g4.5 comes from this	
4483	7511539	gene~cDNA EST yk349a4.3 comes from this gene~cD>	6
		gill 11288486 pir  T49552 hypothetical protein	
		B21J21.300 [imported] - Neurospora crassa	
		emb CAB91353.1  (AL355929) hypothetical protein	
4484	11288486	[Neurospora crassa]	9.5
		gi 11120676 ref NP_068546.1  putative envelope	
		polyprotein [DG-75 Murine leukemia virus]	
		gb AAG29094.1 AF221065_2 (AF221065) putative	
4488	11120676	envelope polyprotein [DG-75 Murine leukemia virus]	1.8
4400	2001254	gi 3901274 gb AAC78630.1  (AF077821) inducible	
4489	3901274	nitric oxide synthase; iNOS [Canis familiaris]	3.5
		gi 11595582 emb CAC18184.1  (AL451014) conserved	
4491	11595582	hypothetical protein [Neurospora crassa]	3
		gi 5835447 ref NP 008371.1 ND6 13186 NADH	
		dehydrogenase subunit 6 [Onchocerca volvulus]	
		pir T11066 NADH dehydrogenase (ubiquinone) (EC	
		1.6.5.3) chain 6 - nematode (Onchocerca volvulus)	
		mitochondrion gb AAC61613.1  (AF015193) NADH	
4493	5835447		1 -
		gi 13814848 gb AAK41824.1  (AE006774) SSV1	
		hypothetical 14.8 kd protein (orf B-129) homolog	_
4494	13814848	[Sulfolobus solfataricus]	5.4
		gi 14529727 cmb CAC42176.1  (AL135758)	
		dM117J5.1 (novel protein similar to anonymous human,	
4500	14529727	fly, worm and yeast proteins) [Mus musculus]	7E-97

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 13512594 gb AAK28688.1  (AF078553) unknown	
4504	13512594	function U3 [Ehrlichia canis]	8.8
		gi 6635084 cmb CAB64573.1  (AL135930) hypothetical	
4506	6635084	protein L4738.02 [Leishmania major]	3.2
		gi 11467470 ref NP_043616.1  50S ribosomal protein	
		L4 [Odontella sinensis] sp P49546 RK4_ODOSI	
		CHLOROPLAST 50S RIBOSOMAL PROTEIN L4	
		pir  S78275 ribosomal protein L4, chloroplast -	
		Odontella sinensis chloroplast emb CAA91648.1	
4508	11467470	(Z67753) 50S ribosomal protein L4 [Odontella sinensis]	5.9
		gi 4512681 gb AAD21735.1  (AC006931) hypothetical	
4509	4512681	protein [Arabidopsis thaliana]	0.73
		gi 5835704 ref NP 008519.1 ND4 15045 NADH	
		dehydrogenase subunit 4 [Rhipicephalus sanguineus]	
		sp O99825 NU4M_RHISA NADH-UBIQUINONE	
		OXIDOREDUCTASE CHAIN 4 pir T11162 NADH	
		dehydrogenase (ubiquinone) (EC 1.6.5.3) chain 4	
		[similarity] - hardbacked tick (Rhipicephalus	
		sanguineus) mitochondrion gb AAD05526.1	
		(AF081829) NADH dehydrogenase 4 [Rhipicephalus	
4510	5835704	sanguineus]	6.8
		gi 8922500 ref NP 060600.1  hypothetical protein	
		FLJ10539 [Homo sapiens] dbj BAA91669.1	
4511	8922500	(AK001401) unnamed protein product [Homo sapiens]	2E-13
1011	0722300	(Factor For Jamanica protein product [Figure sapiens]	2E-13
		gi 12018296 ref NP 072138.1  CDC10 (cell division	
		cycle 10, S.cerevisiae, homolog) [Rattus norvegicus]	
		spiO9WVC0iSEP7 RAT SEPTIN 7 (CDC10	
		PROTEIN HOMOLOG) gblAAD37861,1 AF142759 1	
4514	12018296	(AF142759) CDC10 [Rattus norvegicus]	6.9
		gi 11357368 pir  T48358 hypothetical protein	
		F12E4.100 - Arabidopsis thaliana emb[CAB83293.1]	
4518	11357368	(AL162751) putative protein [Arabidopsis thaliana]	1.1
		gi 6754038 ref[NP_034456.1  glycoprotein 1a, alpha	
4520	(75 4020	polypeptide [Mus musculus] gb AAC53320.1  (U91967)	0.50
4320	0/34038	platelet glycoprotein Ib-alpha [Mus musculus]	0.53

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ ID NO	ACCESS				
NO	N	DESCRIPTION	P VALUE		
		gi 7437434 pir  D72038 DNA topoisomerase I CP1103			
		[imported] - Chlamydophila pneumoniae (strains	}		
		CWL029 and AR39) gb AAD18907.1  (AE001658)			
		DNA Topoisomerase I-Fused to SWI Domain			
		[Chlamydophila pneumoniae CWL029]			
		gb AAF38871.1  (AE002266) DNA topoisomerase I			
		[Chlamydophila pneumoniae AR39] dbj BAA98977.1			
		(AP002547) DNA topoisomerase I-fused to SWI			
4521	7437434	domain [Chlamydophila pneumoniae J138]	8.2		
ĺ l		gi 1514669 emb CAA87082.1  (Z46958) adenyl cyclase			
4525	1514669	[Xenopus laevis]	0.86		
İ					
		gi 6321382 ref NP_011459.1  similar to S. pombe sds23;			
		Sds23p [Saccharomyces cerevisiae]			
		sp[P53172[YGF6_YEAST HYPOTHETICAL 58.1 KD			
		PROTEIN IN UBC2-OLE1 INTERGENIC REGION			
		pir  S64060 probable membrane protein YGL056c -			
		yeast (Saccharomyces cerevisiae) emb CAA96759.1			
4526	6321382		4.2		
		gi 7487690 pir  T01961 hypothetical protein T5H22.5 -			
		Arabidopsis thaliana gb AAC62796.1  (AF096372)			
		contains similarity to reverse transcriptase (Pfam:			
		PF00078 rvt, E=4.3e-08) [Arabidopsis thaliana]			
		emb CAB80803.1  (AL161498) similarity to			
4528	7487690	[Arabidopsis thaliana]	3.9		
4500	10000000	gi 12839203 dbj BAB24467.1  (AK006222) putative			
4530	12839203	[Mus musculus]	9.2		
4501	110000	gi 1177607 emb CAA63219.1  (X92485) pva1			
4531	1177607	[Plasmodium vivax]	0,11		
		gi 7506781 pir  T24250 hypothetical protein R53.3a -			
		Caenorhabditis elegans emb CAA91353.1  (Z66515)			
		contains similarity to Pfam domain: PF00096 (Zinc			
4534	75 <b>067</b> 81	finger, C2H2 type), Score=105.7, E-value=2.9e-28,			
4334	/506/81	N=5 [Caenorhabditis elegans]	3.2		
4520	14041627	gi 14041637 cmb CAC38421.1  (AJ302647) POL			
4538	14041637	protein [Human immunodeficiency virus type 1]	2.8		

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r		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Desteins)
SEQ ID	ACCESS	Table 3B Rearest Reguloof (Blastix Vs. 1401-Redundant	riotenis)
NO	N	DESCRIPTION	P VALUE
		gi 6324843 ref NP_014912.1  Required for viability in	
		the absence of Cin8p; Pac1p [Saccharomyces	
		cerevisiae] sp[P39946[PAC1_YEAST PAC1 PROTEIN	
		pir  S67166 PAC1 protein - yeast (Saccharomyces	
		cerevisiae) emb CAA61775.1  (X89633) hypothetical	
		protein [Saccharomyces cerevisiae] gb AAB00685.1	
		(U16827) Pac1p [Saccharomyces cerevisiae]	
		emb CAA99493.1  (Z75177) ORF YOR269w	
4546	6324843	[Saccharomyces cerevisiae]	4.2
İ		gi 14424433 ref[NP_077816.1  ATPase, Class V, type	
		10C; ATPase type IV, phospholipid transporting (P-	
		type) (putative) [Homo sapiens] gb[AAK33100.1]	
	ļ	(AY029504) aminophospholipid-transporting ATPase	
		[Homo sapiens] dbi BAB47392.1  (AB051358) putative	
4555	14424433	aminophospholipid translocase [Homo sapiens]	3E-10
		gi 6714740 cmb CAB66205.1  (AL136502) putative	
4560	6714740	prolyl aminopeptidase. [Streptomyces coelicolor A3(2)]	4.7
		gi 6724176 gb AAF26878.1 AF196232_1 (AF196232)	
4562	6724176		7.4
		gi 14783807 ref XP 027541.1  hypothetical protein	
4563	14783807	FLJ21858 [Homo sapiens]	3.1
		gi 6478266 gb AAF13781.1 AF129403_2 (AF129403)	
		gamma subunit of membrane-bound ATP synthase	
4564	6478266	[Buchnera aphidicola]	2
		HTTOORIGE HTTOCOCIE	
		gi 7509812 pir  T26861 hypothetical protein Y43F8B.5 - Caenorhabditis elegans emb CAA21513.1  (AL032623)	
		contains similarity to Pfam domain: PF00188 (SCP-like	
		extracellular protein), Score=34.9, E-value=4.2e-09.	
4566	7509812	N=2 [Caenorhabditis elegans]	5.7
		gi 5230656 gb AAD40953.1 AF148934_1 (AF148934)	
4567	5230656	phantastica [Lycopersicon esculentum]	7.9
4570	1453030	gi 14732381 ref XP_050222.1  hypothetical protein	0.0000
4578	14732381	XP_050222 [Homo sapiens]	0.0009
4501	1/2/01/1	gi 14349161 dbj BAB60707.1  (AB049622) ficolin 4	2.5
4581	14349161	[Halocynthia roretzi] gi 3599476 gb AAC69336.1  (AF084637) serendinity	2.5
4585	3500476	gij3599476 gb AAC69336.1  (AP084637) serendinity  alpha protein [Drosophila virilis]	2.4
4000	JJJ774/0	larbae brotem ID1020bing Annis l	4.4

WO 02/14500

PCT/US01/25840

	г	Table 2D March 19 Color 27 Day	
SEO ID		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
NO NO	ACCESS	DEGGDYDWGAY	
NU	N	DESCRIPTION	P VALUE
4586	145510	gi 145512 gb AAC41418.1  (M55661) colonisation	
4380	145512	factor antigen d' [Escherichia coli]	8.9
		gi 7499104 pir  T20906 hypothetical protein F14F7.1 -	
	1	Caenorhabditis elegans emb CAB04111.1  (Z81503)	
		predicted using Genefinder~contains similarity to Pfam	
		domain: PF01484 (Nematode cuticle collagen N-	
	<b>!</b>	terminal domain), Score=31.5, E-value=6.2e-06,	
		N=1~cDNA EST yk56g5.3 comes from this	
4500		gene~cDNA EST yk56g5.5 comes from this gene	
4590	7499104	[Caenorhabditi>	9.8
		gi 7267234 emb CAB80841.1  (AL161501) putative	
4591	7267234	sugar transporter [Arabidopsis thaliana]	9.9
	ł	gi 401050 sp Q01656 RSP4_CHLRE FLAGELLAR	
		RADIAL SPOKE PROTEIN 4 pir  A44498 radial spoke	
	}	protein 4 - Chlamydomonas reinhardtii gb AAA33092.1	
		(M87526) flagellar radial spoke protein	
4593	401050	[Chlamydomonas reinhardtii]	0.31
4-04		gi 7300719 gb AAF55865.1  (AE003734) CG5862 gene	
4594	7300719	product [Drosophila melanogaster]	1.5
4506		gi 12851516 dbj BAB29072.1  (AK013941) putative	
4596	12851516	[Mus musculus]	1.1
		gi 14193314 gb AAK55896.1 AF267213_2 (AF267213)	
4500		ATP synthase gamma subunit [Candidatus Carsonella	
4598	14193314		5.2
		gi 13898998 gb AAK48930.1 AF359251_1 (AF359251)	
4500	12000000	extracellular polypeptide Ecp76 [Chlamydomonas	
4599	13898998	reinhardtii]	0.17
		Topography man occasionally	
		gi 8922500 ref[NP_060600.1  hypothetical protein	
4601	0000500	FLJ10539 [Homo sapiens] dbj[BAA91669.1]	
4001	8922500	(AK001401) unnamed protein product [Homo sapiens]	4.7
		gi 7494152 pir  T18410 carbamoyl-phosphate synthase	
		(glutamine-hydrolyzing) (EC 6.3.5.5) II - malaria	
		parasite (Plasmodium falciparum) gb AAA29522.1	
4605	7494152	(L32150) carbamoyl phosphate synthetase II [Plasmodium falciparum]	0.7
4000	7494152	gi 6178092 dbj BAA86168.1  (AB031705) ORF2	8.7
4607	6178092	gi 01/8092 d0] BAA86168.1  (AB031/05) ORF2 protein [TT virus]	2.4
4007	01/8092	protein [11 vitus]	3.4
		gi 122078 sp P02301 H34 MOUSE HISTONE H3.4	
		(EMBRYONIC) emb CAA24131.1  (V00754) reading	
4608	122078	frame histone H3 [Mus musculus]	0.059
7000	122010	Trans musculas	0.059

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	Treatment (Billion (Billion 18, 116))	T Totalis)
NO	N	DESCRIPTION	P VALUE
		gi 2598224 emb CAA69955.1  (Y08696) aldehyde	- 11202
4610	2598224	dehydrogenase [Gluconacetobacter europaeus]	2.1
		, , , , , , , , , , , , , , , , , , , ,	
		gi 12382242 gb AAG53080.1 AF263824 1 (AF263824)	
4612	12382242		3.2
1012	12502212	5 rector rotated protein [Ecisimania donovani]	3.2
		-31 40522 (JB 3D 4 4 120 (0 11 (D0 (0 40) 1	
		gi 1405336 dbj BAA13060.1  (D86240) hypothethecal membrane transporter [Staphylococcus aureus]	
		gb AAD21958.1  (AF101234) putative membrane	
		protein DltB [Staphylococcus aureus] dbj BAB42033.1	
		(AP003132) DltB membrane protein [Staphylococcus	
		aureus subsp. aureus N315] dbj BAB57095.1	
4610	1405006	(AP003360) DItB membrane protein [Staphylococcus	
4613	1405336	aureus subsp. aureus Mu50]	7.4
		gi 6321803 ref[NP_011879.1  PolyA-binding protein;	
		Mip6p [Saccharomyces cerevisiae]	
		sp[P38760]YHH5_YEAST HYPOTHETICAL 75.9	
		KDA PROTEIN IN SPO13-ARG4 INTERGENIC	
		REGION pir S46788 PES4 protein homolog YHR015w	
		- yeast (Saccharomyces cerevisiae) gb AAB68942.1	
4616	6321803	(U10400) Yhr015wp [Saccharomyces cerevisiae]	5.3
		gi 14749154 ref XP_031524.1  AF15q14 protein [Homo	
4618	14749154		9E-94
		gi 12837658 dbj BAB23899.1  (AK005241) putative	
		[Mus musculus] dbj BAB26680.1  (AK010067) putative	
4621	12837658	[Mus musculus]	6.8
	(	gi 9628166 ref[NP_042752.1  CD2 homolog [African	
		swine fever virus] pir  A40678 T-cell adhesion receptor	
		CD2 homolog - African swine fever virus	
		gb AAA42691.1  (L16864) cd2 homologue [African	
		swine fever virus] gb AAA65288.1  (U18466) CD2	
		homolog [African swine fever virus] prf[2113434BJ	
4622	9628166	CD2-like protein [African swine fever virus]	7.6
		gi 56691 emb CAA68549.1  (Y00497) precursor (AA -	
4625	56691	24 to 198) [Rattus norvegicus]	4.2
		gi 10946710 ref NP_067350.1  Rhesus blood group-	
		associated B glycoprotein; Rh type B glycoprotein [Mus	
		musculus] gb AAF19371.1  (AF193808) Rh type B	
4628	10946710	glycoprotein [Mus musculus]	9.9

	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESS	Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total and Total	T Totalis)		
NO	N	DESCRIPTION	P VALUE		
	1	gi 7507118 pir  T24482 hypothetical protein T05A1.2 -			
		Caenorhabditis elegans emb CAA92476.1  (Z68219)			
		contains similarity to Pfam domain: PF01391 (Collagen			
		triple helix repeat (20 copies)), Score=86.4, E-			
		value=1.8e-22, N=2; PF01484 (Nematode cuticle			
		collagen N-terminal domain), Score=23.9, E-			
		value=0.0012, N=1~cDNA EST CEMSE21F comes			
4631	7507118	from this >	3.6		
		gi 2072964 gb AAC51271.1  (U93569) putative p150			
4634	2072964	[Homo sapiens]	0.12		
		gi 13385228 ref NP 080036.1  RIKEN cDNA			
		4933428I03 gene [Mus musculus] dbi BAB30529.1			
		(AK016968) putative [Mus musculus] dbj BAB30656.1			
4636	13385228	(AK017257) putative [Mus musculus]	2.1		
		gi 13637800 ref XP_017899.1  hypothetical protein			
4639	13637800	FLJ12673 [Homo sapiens]	2E-11		
		gi 14759884 ref XP 010198.3  64448 [Homo sapiens]			
		sp Q9BZS1 FXP3 HUMAN FORKHEAD BOX			
		PROTEIN P3 (ZINC FINGER PROTEIN JM2)			
		(SCURFIN) gb AAG53607.1 AF277993 1 (AF277993)			
4645	14759884	scurfin [Homo sapiens]	0.000002		
		gi 9758192 dbj BAB08666.1  (AB018109)			
4651	9758192	pectinesterase [Arabidopsis thaliana]	5.5		
		gi 9989055 gb AAG10818.1 AC011808_6 (AC011808)			
4653	9989055	Hypothetical protein [Arabidopsis thaliana]	6E-11		
		gi 560700 gb AAB31458.1  gibbon ape leukemia virus			
		receptor [Mus musculus=Japanese feral mice, spp.			
4658	560700	molossinus, susceptible cells, Peptide, 680 aa]	7		
		gi 3878238 emb CAA81588.1  (Z27078) cDNA EST			
		yk181g1.5 comes from this gene~cDNA EST			
		yk153a11.5 comes from this gene~cDNA EST			
		yk465b1.5 comes from this gene [Caenorhabditis			
4659	3878238	elegans]	1.3		

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	+ Destaine)
SEQ ID	ACCESS	Tuble SB Treatest Treignoor (Blasta Vs. 14011-Rediffically	i Fiotenis)
NO	N	DESCRIPTION	P VALUE
		gi 141028 sp P04540 NU5M_TRYBB NADH-	
	!	UBIQUINONE OXIDOREDUCTASE CHAIN 5	
	i	pir  QQUTC5 NADH dehydrogenase (ubiquinone) (EC	ł
		1.6.5.3) chain 5 - Trypanosoma brucei mitochondrion	
		gb AAB59225.1  (M14820) NADH dehydrogenase	]
		subunit 5 [Trypanosoma brucei] emb[CAB57807.1]	
		(X01094) unidentified reading frame 10 [Trypanosoma	
4663	141028	brucei]	0.54
		gi 5442110 gb AAD43259.1 AF126468_2 (AF126468)	
4664	5442110	protease [Simian retrovirus type 2]	0.3
		gi 12860471 dbj BAB31968.1  (AK020018) putative	
4666	12860471	[Mus musculus]	7.6
		gi 4103974 gb AAD05047.1  (AF030414) FeMo protein	
	ĺ	of nitrogenase alpha subunit; NifD [Gluconacetobacter	
4669	4103974		9.1
		gi 2072953 gb AAC51264.1  (U93565) putative p150	
4678	2072953	[Homo sapiens]	0.057
		gi 7208454 gb AAF40208.1 AF233885_1 (AF233885)	
4679	7208454	phospholipase C-like protein [Mus musculus]	9.6
		gi 14193286 gb AAK55875.1 AF267206_2 (AF267206)	
		ATP synthase gamma subunit [Candidatus Carsonella	
4680	14193286		9
		gi 11361549 pir  A82772 hypothetical protein XF0722	
		[imported] - Xylella fastidiosa (strain 9a5c)	
		gb AAF83532.1 AE003914_13 (AE003914)	
4681	11361549	hypothetical protein [Xylella fastidiosa 9a5c]	3.1
		gi 7463298 pir  A70144 hypothetical protein BB0354 -	
[		Lyme disease spirochete gb AAC66737.1  (AE001141)	
		B. burgdorferi predicted coding region BB0354	
4682	7463298	[Borrelia burgdorferi]	5.4
		gi 9631610 ref NP 048389.1  contains Pro-rich Px	
		motif, PAPK (8X); similar to Thermoproteus virus	
		protein TPX, corresponds to Swiss-Prot Accession	
		Number P19275 [Paramecium bursaria Chlorella virus	
		1] pir  T17531 proline-rich protein A41R - Chlorella	
		virus PBCV-1 gb AAC96409.1  (U42580) contains Pro-	
	'	rich Px motif, PAPK (8X); similar to Thermoproteus	
		virus protein TPX, corresponds to Swiss-Prot Accession	
		Number P19275 [Paramecium bursaria Chlorella virus	
4685	9631610	1]	1.8

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	I	Table 3B Nearest Neighbor (BlastX vs. Non-Redundam	Proteins)
SEQ ID	ACCESS	See Comment	T Totolia)
NO	_ N	DESCRIPTION	P VALUE
		gi 14754644 ref XP_030896.1  similar to hypothetical	
4688	14754644	protein MGC5384 (H. sapiens) [Homo sapiens]	9E-48
		gi 2865163 dbj BAA24799.1  (AB007836) Hic-5 [Homo	
4689	2865163	sapiens]	2E-16
		gi 6760239 gb AAF28263.1 AF178655_4 (AF178655)	
4693	6760239	fusion F [Bovine parainfluenza virus 3]	9.8
		gi 13446199 cmb CAC34985.1  (AL035536) very	
4698	13446199	hypothetical protein [Schizosaccharomyces pombe]	5.2
		gi 6320491 ref NP_010571.1  synaptonemal complex	
		protein; Zip1p [Saccharomyces cerevisiae]	
		sp P31111 ZIP1_YEAST SYNAPTONEMAL	
		COMPLEX PROTEIN ZIP1 pir S70115 ZIP1 protein -	
		yeast (Saccharomyces cerevisiae) gb AAB64474.1	
		(U51031) Zip1p: Synaptonemal complex protein (Swiss	
		Prot. accession number P31111). [Saccharomyces	
4699	6320491	cerevisiae]	0.001
		gi 416704 sp Q03376 BAR3_CHITE BALBIANI RING	
		PROTEIN 3 PRECURSOR pir  S08167 Balbiani ring 3	
		protein - midge (Chironomus tentans) emb CAA36506.1	
4703	416704	(X52263) balbiani ring 3 (BR3) [Chironomus tentans]	5.4
		gi 6005948 ref NP_009118.1  WW domain-containing	
		binding protein 4; formin binding protein 21 [Homo	
		sapiens] ref[XP_007153.1  WW domain-containing	
		binding protein 4 [Homo sapiens] ref[XP_049374.1]	
		WW domain-containing binding protein 4 [Homo	
		sapiens] ref XP_049375.1  WW domain-containing	
		binding protein 4 [Homo sapiens] gb AAC34811.1	
4709		(AF071185) formin binding protein 21 [Homo sapiens]	0.007
		gi 629166 pir  S39791 neurotoxin - Clostridium	
4713		botulinum	2.5
		gi 4096360 gb AAC99858.1  (U31159) CR16 [Rattus	
		norvegicus] gb AAC99859.1  (U31169) SH3 domain	
4719		binding protein [Rattus norvegicus]	4.3
		gi 11352438 pir  F83161 pyocin protein PA3866	
		[imported] - Pseudomonas aeruginosa (strain PAO1)	
		gb AAG07253.1 AE004804_1 (AE004804) pyocin	
4723		protein [Pseudomonas aeruginosa]	9.1
		gi 14761847 ref XP_017198.2  hypothetical protein	
4725	14761847	FLJ12085 [Homo sapiens]	0.0007

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
110		gi 7475497 pir  B70045 hypothetical protein yvpB -	PVALUE
		Bacillus subtilis gb AAC67292,1  (AF017113) YvpB	
		[Bacillus subtilis] emb CAB15499.1  (Z99121) yvpB	
4726	7475497	[Bacillus subtilis] emo[CAB13499.1] (Z99121) yvpB	9
4720	1413431	[Dacinus subulis]	9
		gi 11466185 ref NP 066508.1  NADH dehydrogenase	
		subunit 11 [Naegleria gruberi]	
		gb AAG17786.1 AF288092 11 (AF288092) NADH	
4731	11/66195	dehydrogenase subunit 11 [Naegleria gruberi]	6.2
4/31	11400183	gi 9802589 gb AAF99791.1 AC012463 8 (AC012463)	0.2
4735	9802589	T2E6.16 [Arabidopsis thaliana]	7.7
4733	9002309	gi 7512245 pir T03849 Fas-binding protein Daxx -	1.1
		green monkey gb AAB66586.1  (AF015957) Fas-	
4741	7512245	binding protein Daxx [Cercopithecus aethiops]	7.1
4/41	7312243	gi 345660 pir  A45031 cysteine-rich fibroblast growth	7.1
		factor receptor - chicken gb AAA48769.1  (M95766)	
		cysteine-rich fibroblast growth factor receptor [Gallus	
4746	345660	gallus]	1
17.10	3 13 000	gunusj	1
		gi 6324550 ref NP 014619.1  mitochondrial initiation	
		factor 2; Ifm1p [Saccharomyces cerevisiae]	
		sp P25038 IF2M YEAST TRANSLATION	
		INITIATION FACTOR IF-2, MITOCHONDRIAL	
		PRECURSOR (IF-2MT) (IF-2(MT)) pir S66706	
		translation initiation factor IF-2, mitochondrial - yeast	
		(Saccharomyces cerevisiae) emb CAA99023.1	
4750	6324550	(Z74765) ORF YOL023w [Saccharomyces cerevisiae]	8.1
		gi 12249161 ref NP 066211.2  cytochrome c oxidase	
		subunit III [Schistosoma mansoni] gb[AAG13163.2]	
		(AF216698) cytochrome c oxidase subunit 3	
4751	12249161	[Schistosoma mansoni]	6.6
		gi 10835218 ref NP 004609.1  topoisomerase (DNA) III	
		alpha [Homo sapiens] ref[XP 008635.1] topoisomerase	
		(DNA) III alpha [Homo sapiens]	
		sp Q13472 TP3A HUMAN DNA TOPOISOMERASE	
		III ALPHA gb AAB03694.1  (U43431) DNA	
4758	10835218	topoisomerase III [Homo sapiens]	9.4
		gi 9294528 dbj BAB02791.1  (AB024034)	
		gb AAF50915.1~gene_id:MDC11.5~similar to unknown	
4759	9294528	protein [Arabidopsis thaliana]	7.1
		gi 6624755 cmb CAB63872.1  (AJ251846) OTX5b	
4760	6624755	protein [Xenopus laevis]	8.8

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundam	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 14768465 ref XP_038469.1  hypothetical protein	
4762	14768465	XP_038469 [Homo sapiens]	0.88
		gi 7290081 gb AAF45547.1  (AE003418) fz3 gene	
4768	7290081	product [alt 2] [Drosophila melanogaster]	4.7
4770	12259052	gijl 3358052[ref[NP_078326.1] unique hypothetical membrane lipoprotein [Ureaplasma urealyticum] pir[H82883 hypothetical protein UU489 [imported] - Ureaplasma urealyticum gb]AAF30901.1[AE002147_2 (AE002147) unique hypothetical membrane lipoprotein [Ureaplasma urealyticum]	
4770	15556052		0.95
4774	12698025	gi 12698025 dbj BAB21831.1  (AB051527) KIAA1740 protein [Homo sapiens]	0.24
4//4	12098023	protein [riomo sapiens]	0.24
4784	116910	gi 116910 sp P10169 CON8_NEUCR CONIDIATION- SPECIFIC PROTEIN 8 pir  S02210 con-8 protein - Neurospora crassa emb CAA30092.1  (X07040) con-8 [Neurospora crassa]	4.3
4704	110710	gi 3694664 gb AAC62434.1  (AC004893) similar to	4.3
4788	3694664	NEDD-4 (KIA0093); similar to P46934 (PID:g1171682) [Homo sapiens]	8.2
4790	1731439	gi 1731439 sp P51505 ZN80_MACMU ZINC FINGER PROTEIN 80 emb CAA61771.1  (X89629) znf80 [Macaca mulatta]	1.3
4794	11257183	gi 11257183 pir F82319 transketolase 1 VC0473 [imported] - Vibrio cholerae (group O1 strain N16961) gb AAF93646.1  (AE004133) transketolase 1 [Vibrio	9.1
7774	1120/103		7,1
		gil1146520[nrf]NP_044769.1] SecY-type transporter protein [Reclinomonas americana] piri[S78151 secY protein homolog - Reclinomonas americana (ATCC 50394) mitochondrion gb]AD11884.1] (AF007261) SecY-type transporter protein [Reclinomonas	
4795	11466520	americana]	2.3
4799	5869811	gi 5869811 emb CAB55552.1  (AJ243538) Fox2 protein [Glomus mosseae]	8.5

WO 02/14500

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 6324187 ref NP_014257.1  Ammonia transport	
		protein; Mep2p [Saccharomyces cerevisiae]	
		sp P41948 MEP2_YEAST AMMONIUM	
		TRANSPORTER MEP2 pir  S51089 ammonium	
		transport protein MEP2 - yeast (Saccharomyces	
		cerevisiae) emb CAA58587.1  (X83608) ammonium	
		transporter [Saccharomyces cerevisiae]	
		emb CAA86884.1  (Z46843) NH3 permease	
		[Saccharomyces cerevisiae] emb CAA96025,1	
4801	6324187	(Z71418) ORF YNL142w [Saccharomyces cerevisiae]	8.3
		gi 14738182 ref XP_038231.1  KIAA0793 gene product	
4805		[Homo sapiens]	3E-16
4806	481861	gi 481861 pir  S39796 aggrecan precursor - chicken	0.065
		gi 462116 sp Q05394 FMLR_RABIT FMET-LEU-PHE	
		RECEPTOR (FMLP RECEPTOR) (N-FORMYL	
		PEPTIDE RECEPTOR) (FPR) (N-FORMYLPEPTIDE	
		CHEMOATTRACTANT RECEPTOR) pir  A46520 N-	
		formyl peptide receptor - rabbit gb AAA31254.1	
		(M94549) N-formyl peptide receptor [Oryctolagus	
4808	462116	cuniculus]	2.2
		gi 12832739 dbj BAB22236.1  (AK002621) putative	
4809	12832739	[Mus musculus]	1.1
		gi 464359 sp P33295 PEPC_ASPNG SUBTILISIN-	
- 1		LIKE SERINE PROTEASE PEPC PRECURSOR	
		pir  JU0146 serine proteinase (EC 3.4.21) precursor -	
4813	161050	Aspergillus niger gb AAA32702.1  (M96758) serine	
4813	464359	protease [Aspergillus niger]	0.91
4814	14741406	gi 14741436 ref XP_032168.1  hypothetical protein XP_032168 [Homo sapicns]	
4614	14/41436	XP_032168 [Homo sapiens]	1.3
J		gi 6322061 ref NP_012136.1  Yil130wp	
		[Saccharomyces cerevisiae] sp P40467 YIN0_YEAST	
		PUTATIVE 108.8 KD TRANSCRIPTIONAL	
		REGULATORY PROTEIN IN FKH1-STH1	
- 1		INTERGENIC REGION pir  S48404 probable	
		membrane protein YIL130w - yeast (Saccharomyces	
		cerevisiae) emb CAA86148.1  (Z38059) orf, len: 964, CAI: 0.15, possible regulatory protein [Saccharomyces	
		CAL U.13, DOSSIDIC REQUISIORY Drofein I Saccharomyces	
4821	6322061		67
4821	6322061	cerevisiae]	6.7

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Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESS			
NO	N	DESCRIPTION	P VALUE	
		gi 13926079 gb AAK49526.1 AF356082_1 (AF356082)		
	1	C2H2 zinc finger transcription factor JING [Drosophila		
4825	13926079	melanogaster]	2.7	
		gi 1209370 gb AAA91081.1  (L33971) EBV BKRF4		
4827	1209370	homologue [Leporid herpesvirus 1]	8	
	ĺ	gi 11359537 pir  T51023 hypothetical protein B7F21.40		
		[imported] - Neurospora crassa emb CAB97476.1		
		(AL389901) conserved hypothetical protein		
4829	11359537	[Neurospora crassa]	0.57	
	l			
	1	gi 13629392 sp P97523 MET_RAT HEPATOCYTE		
		GROWTH FACTOR RECEPTOR PRECURSOR		
		(MET PROTO-ONCOGENE TYROSINE KINASE)		
		(C-MET) (HGF RECEPTOR) (HGF-SF RECEPTOR)		
4832	13629392	emb CAA65582.1  (X96786) c-met [Rattus norvegicus]	1.3	
	ľ	gi 4508019 ref NP_003449.1  bassoon (presynaptic		
		cytomatrix protein); neuronal double zinc finger protein;		
		zinc finger protein 231 [Homo sapiens] gb AAC83555.1		
		(AF052224) neuronal double zinc finger protein [Homo		
4834	4508019	sapiens]	1.5	
		gi 5734374 emb CAB52680.1  (AJ133723) Ran-binding		
4835	5734374	protein 2 [Bos taurus]	4.7	
		gi 3281932 emb CAA76796.1  (Y17584) beta-lactamase		
		class A [Escherichia coli] emb CAC43180.1		
		(AJ277416) TEM-29 ES-beta-lactamase [Escherichia		
4837	3281932	coli]	9.4	
		gi 15021417 gb AAK77694.1 AF369029_25		
		(AF369029) ORF25, gene family 4 [white spot		
4839	15021417	syndrome virus]	9.4	
		gi 7507283 pir  T32024 hypothetical protein T06D4.4 -		
		Caenorhabditis elegans gb AAB66123.1  (AF016673)		
		similar to endothelin-converting enzymes		
4841	7507283	[Caenorhabditis elegans]	6.7	
		gi 1165132 emb CAA64491.1  (X95193) homeobox-		
4843	1165132	leucine zipper protein [Pimpinella brachycarpa]	8.5	

NO   N   DESCRIPTION   P VALUE			Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
gij 11466755 refiNP_039351.1  ORF191 [Marchantia polymorpha] splP06266 NU6C_MARPO NADH-PLASTOQUINONE OXIDOREDUCTASE CHAIN 6, CHI.OROPLAST pir  DEL/NN NADH dehydrogenase (ubiquinone) (EC 1.6.5.3) chain 6 - liverwort (Marchantia polymorpha) chloroplast emb CAA28137.1  (X04465) ORF191 [Marchantia emb CAA28137.1  (X04465) ORF191 [Marchantia emb CAA28137.1  (X04465) ORF191 [Marchantia emb CAA28137.1  (X04465) ORF191 [Marchantia emb CAA28137.1  (X04465) ORF191 [Marchantia emb CAA28137.1  (X04465) ORF191 [Marchantia emb CAA28137.1  (X04465) ORF191 [Marchantia emb CAA28137.1  (X04465) ORF191 [Marchantia emb CAA28137.1  (X04465) ORF191 [Marchantia emb CA2838.1  similar to cleft lip and palate associated transmembrane protein I (H. sapiens) [Homo sapiens] eji 147785 refiXP_04838.1  similar to hypothetical protein FLJ13166 (H. sapiens) [Homo sapiens] eji 1994373 dbj BAB02332.1  (AB019229) gene_id-MDC16.14-unknown protein [Arabidopsis emi]1994373 (thaliana] 4.7  4866 11994373 (thaliana) 4.7  4872 12514590 protein [Escherichia coli O157:H7 ED1933] dbj BAB34856.1  (AP002555) hypothetical protein FLD1933] dbj BAB34856.1  (AP002555) hypothetical protein [Escherichia coli O157:H7 ED1933] dbj BAB34856.1  (AB058718) KIAA1815 efinal fill of the protein [Escherichia coli O157:H7 ED1933] dbj BAB34856.1  (AB058718) KIAA1815 efinal fill of the protein [Homo sapiens] eji 7506698[pir T24214 hypothetical protein R134.2 - Caenorhabditis elegans emb CAB05777.1  (Z83236) contains similarity to Pfam domain: PF00083 (Sugar (and other) transporter), Scores—89.7, E-value—00037, N=-cDNA EST yk653al.1.3 comes from this gene-cDNA EST yk653al.1.5 comes from this gene-cDNA EST yk653al.1.5 comes from this gene-cDNA EST yk653al.1.5 comes from this gene-cDNA EST yk653al.1.5 comes from this gene-cDNA EST yk653al.1.5 comes from this gene-cDNA EST yk653al.1.5 comes from this gene-cDNA EST yk653al.1.5 comes from this gene-cDNA EST yk653al.1.5 comes from this gene-cDNA EST yk653al.1.5 comes from this gene-cDNA EST yk653al.1.5 comes from this gene-cDNA EST	SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
Dolymorpha  sp P06266 NU6C_MARRO NADH-PLASTOQUINONE OXIDDREDUCTASE CHAIN 6, CHLOROPLAST prig DELVN6 NADH dehydrogenase (ubiquinone) (EC 1.6.5.3) chain 6 - liverwort (Marchantia polymorpha) chloroplast emb CAA28137.1  (X04465) ORF191 [Marchantia polymorpha]				1 11202
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(ubiquinone) (EC 1.6.5.3) chain 6 - liverwort (Marchantia polymorpha) chloroplast emb[CAA2813-1] (LV04465) ORF191 [Marchantia emb[CAA2813-1] (LV04465) ORF191 [Marchantia emb[CAA2813-1] (LV04465) ORF191 [Marchantia emb[CAA2813-1] (LV04465) ORF191 [Marchantia emb[CAA2813-1] (LV04465) ORF191 [Marchantia emb[CAA2813-1] (LV04465) ORF191 [Marchantia emb[CAA2813-1] (LV04465) ORF191 [Marchantia emb[CAA2813-1] (LV04465) ORF191 [Marchantia emb[CAB291] (AF363973) (AF363973) employed (AF363973) (AF363973) (AF363973) (AF363973) employed (AF36978) (AF36978) (AF36978) (AF36978) (AF36978) (AF36978) (AF36978) (AF36978) (AF36978) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF36978788) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF3697878) (AF36978878) (AF3697878) (AF3697878) (AF36978878) (AF3697878) (AF36978878) (AF36978878) (AF36978878) (AF36978878) (AF3697878) (AF36978878) (AF36978878) (AF36978878) (AF36978878) (AF36978878) (AF3697887			PLASTOQUINONE OXIDOREDUCTASE CHAIN 6,	
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emb CAA28137.1  (X04465) ORF191 [Marchantia   9.5     emb CAA28137.1  (X04465) ORF191 [Marchantia   9.5     gill4010605] polymorpha  9.5     gill4010605] semaphorin 6C [Mus muscalus] 6.3     gill4755906 reifXP_044358.1  similar to cleft lip and palate associated transmembrane protein 1 (H. sapiens)     4853   14755906  [Homo sapiens]   2.7     gill3647785 reifXP_018036.1  hypothetical protein   FIJ13166 [Homo sapiens]   13647785   13166 [Homo sapiens]   1610     4859   13647785   13947785 reifXP_018036.1  hypothetical protein   FIJ13166 [H. sapiens) [Homo sapiens]   6.1     gill 1994373 dbj BAB02332.1  (AB019229)   gene_id:MDC16.14-unknown protein [Arabidopsis   4.7     4866   11994373   4181481   4.7     gill2514590]gblAAG58801.1 AE005316_6 (AE005316)   67f. hypothetical protein [Escherichia coli 0157:H7     EDL933  dbj BAB34856.1  (AP002555) hypothetical   6     gill4017847 dbj BAB47444.1  (AB058718) KIAA1815   7506698   7506698   7506698   7506698   7506698   7506698   750760  77.3   77.3   77.3     4878   7506698   750750 pir[T23590 hypothetical protein R134.2 - 7.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.3   77.				
14845   11466755   polymorpha    9.5	ļ			
14010605 gb]AAK52051.1]AF363973   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF363973)   (AF36397		l		
14010605	4845	11466755		9.5
gij 14755906 rcfiXP_044358.1  similar to cleft lip and palate associated transmembrane protein 1 (H. sapiens)   2.7				
A853   14755906	4849	14010605		6.3
4853   14755906   Homo sapiens   2.7				
gij13647785 reijXP_018036.1  hypothetical protein   FLJ13166 [Homo sapiens] refXP_051378.1] similar to hypothetical protein FLJ13166 [Homo sapiens] refXP_051378.1] similar to hypothetical protein FLJ13166 (H. sapiens) [Homo sapiens]   6.1   gij11994373 dbjjBAB02332.1  (AB019229)   6.1   4.7   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1   6.1	4052	14755006		2.7
FLJ13166 [Homo sapiens] refXP_051378.1] similar to hypothetical protein FLJ13166 [H. sapiens) [Homo of the protein FLJ13166 [H. sapiens)]   Flow of the protein FLJ13166 [H. sapiens)]   Flow of the protein FLJ13166 [H. sapiens)]   Flow of the protein FLS1316 [Homo of the protein FLS1316 [Homo of the protein FLS1316]   Flow of the protein [Arabidopsis]   4.7	4033	14733900		2.1
hypothetical protein FIJ13166 (H. sapiens) [Homo sapiens]	İ			
4859   13647785   sapiens				
gij11994373 tbjjBAB02332.1  (AB019229)   gene_id:MDC16.14-unknown protein [Arabidopsis     4.7     1994373   thaliana    4.7     gij12514590 gb]AAG55801.1 AE005316_6 (AE005316)     orf. hypothetical protein [Escherichia coli O157:H7     EDL933  dbjjBAB34856.1  (AP002555) hypothetical     2514590   protein [Escherichia coli O157:H7     gij14017847 dbjBAB44744.1  (AB058718) KIAA1815     4873   14017847   protein [Homo sapiens]   6E-18     gij7506698 pir1724214 hypothetical protein R134.2 -     4878   7506698   Caenorhabditis elegans   7.3     2875750 pir1723590 hypothetical protein K10H10.1 -     Caenorhabditis elegans emb CAB05777.1  (Z83236)     contains similarity to Pfam domain: PF00083 (Sugar (and other) transporter), Score=89.7, E-value=0 00037,     N=1-cDNA EST yk653a11.3 comes from this     gene-cDNA EST yk653a11.3 comes from this     gene-cDNA EST yk653a11.3 comes from this     gene-cDNA EST yk653a11.3 comes from this     gene-cDNA EST yk653a11.5 comes from this     gene-cDNA EST yk653a11.5 comes from this     gene-cDNA EST yk653a11.5 comes from this     gene-cDNA EST yk653a11.5 comes from this     gene-cDNA EST yk653a11.5 comes from this     gene-cDNA EST yk653a11.5 comes from this     gene-cDNA EST yk653a11.5 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from this     gene-cDNA EST yk653a1.7 comes from thi	4859	13647785		6.1
gene_id:MDC16.14~unknown protein [Arabidopsis   4.7				
4.7	1			
orf, hypothetical protein [Escherichia coli O157:H7   EDL.933] dbjjlBAB34856. [I, (AP002555) hypothetical   EDL.933] dbjjlBAB34856. [I, (AP002555) hypothetical   [Escherichia coli O157:H7] 6   [Escherichia coli O157:H7] 6   [Escherichia coli O157:H7] 6   [Escherichia coli O157:H7] 6   [Escherichia coli O157:H7] 6   [Escherichia coli O157:H7] 6   [Escherichia coli O157:H7] 7   [Escherichia coli O157:H7] 7   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H7] 8   [Escherichia coli O157:H	4866	11994373		4.7
orf, hypothetical protein [Escherichia coli O157:H7]  BDL933 [db]]BAB34856.1 [AP002555] hypothetical  BDL933 [db]]BAB34856.1 [AP002555] hypothetical  gil[14017847][bb]BAB47444.1 [AB058718] KIAA1815  gil[14017847][bb]BAB47444.1 [AB058718] KIAA1815  gil[7505698] pir[721214 hypothetical protein R134.2 -  Caenorhabditis elegans  7.3  gil[7505750][pir[723590 hypothetical protein K10H10.1 - Caenorhabditis elegans emb]CAB05777.1 [(Z83236)  contains similarity to Pfam domain: PF00083 (Sugar  (and other) transporter), Scorre—89.7, E-value=0.00037,  N=1-cDNA EST yk653al1.3 comes from this  gene-cDNA EST yk653al1.3 comes from this  gene-cDNA EST yk653al1.5 comes from this gene  [Caenorhab]  4880 7505750 [Caenorhab]  1.4  gi[2500112][piP71359][RECQ_HAEIN ATP_  DEPENDENT DNA HELICASE RECQ  gb]AAC22387.1 [(032756) ATP-dependent DNA				
EDL933  dbj BAB34856.1  (AP002555) hypothetical			gi 12514590 gb AAG55801.1 AE005316_6 (AE005316)	
4872   12514590   protein [Escherichia coli O157:H7]   6			orf, hypothetical protein [Escherichia coli O157:H7	
14017847   gijl4017847 sbjjBAB47444.1  (AB058718) KIAA1815     4878				
4873   14017847   protein [Homo sapiens]   6E-18	4872	12514590		6
2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007   2007	i			
4878 7506698 Caenorhabditis elegans 7.3  gij7505750[pir][T23590 hypothetical protein K10H10.1 - Caenorhabditis elegans emb]CAB05777.1] (Z83236) contains similarity to Pfam domain: PF00083 Gsugar (and other) transporter), Score=89.7, E-value=0.00037, N=1-cDNA EST yk503h12.5 comes from this gene-cDNA EST yk653al1.3 comes from this gene-cDNA EST yk653al1.5 comes from this gene-CDNA EST yk653al1.5 comes from this gene-CDNA EST yk653al1.5 comes from this gene-CDNA EST yk653al1.5 comes from this gene-CDNA EST yk653al1.5 comes from this gene [1.4]  4880 7505750 [Caenorha> 1.4	4873	14017847		6E-18
gij7505750jpir[T23590 hypothetical protein K10H10.1 - Caenorhabditis elegans emb CAB05777.1  (Z83236) contains similarity to Pfam domain: PF00083 (Sugar (and other) transporter), Score=89.7, E-value=0 00037, N=1=cDNA EST yk503al1.2 comes from this gene=cDNA EST yk503al1.3 comes from this gene=cDNA EST yk503al1.3 comes from this gene gene=cDNA EST yk503al1.3 comes from this gene gene=cDNA EST yk503al1.5 comes from this gene [Caenorha= 1.4  gij2500112 sp P71359 RECQ_HAEIN ATP- DEPENDENT DNA HELICASE RECQ gb AAC22387.1  (U32756) ATP-dependent DNA				_
Caenorhabditis elegans emb CAB05777.1  (Z83236) contains similarity to Pfam domain: PF00083 (Sugar (and other) transporter), Score=89.7, E-value=0 00037, N=1~cDNA EST yk503h12.5 comes from this gene~cDNA EST yk653a11.3 comes from this gene—CDNA EST yk653a11.5 comes from this gene—CDNA EST yk653a11.5 comes from this gene—CDNA EST yk653a11.5 comes from this gene 4880 7505750 [Caenorha> 1.4  gi[2500112]sp[P71359]RECQ_HAEIN ATP— DEPENDENT DNA HELICASE RECQ gb[AAC22387.1] (U32756) ATP-dependent DNA	4878	7506698	Caenorhabditis elegans	7.3
Caenorhabditis elegans emb CAB05777.1  (Z83236) contains similarity to Pfam domain: PF00083 (Sugar (and other) transporter), Score=89.7, E-value=0 00037, N=1~cDNA EST yk503h12.5 comes from this gene~cDNA EST yk653a11.3 comes from this gene—CDNA EST yk653a11.5 comes from this gene—CDNA EST yk653a11.5 comes from this gene—CDNA EST yk653a11.5 comes from this gene 4880 7505750 [Caenorha> 1.4  gi[2500112]sp[P71359]RECQ_HAEIN ATP— DEPENDENT DNA HELICASE RECQ gb[AAC22387.1] (U32756) ATP-dependent DNA	1			
contains similarity to Pfam domain: PF00083 (Sugar (and other) transporter), Score=89.7, E-value=0 00037, N=1-cDNA EST yk503h12.5 comes from this gene-cDNA EST yk653al1.3 comes from this gene-cDNA EST yk653al1.3 comes from this gene 4880 7505750 [Caenorha> 1.4  gi[2500112]sp[P71359]RECQ_HAEIN ATP- DEPENDENT DNA HELICASE RECQ gb[AAC22387.1] (U32756) ATP-dependent DNA				
(and other) transporter), Score—89.7, E-value—0.00037, N=1-cDNA EST yk503h12.5 comes from this gene-cDNA EST yk653al1.3 comes from this gene-cDNA EST yk653al1.5 comes from this gene 4880 7505750 [Caenorha> 1.4  gil2500112 sp P71359 RECQ_HAEIN ATP- DEPENDENT DNA HELICASE RECQ gb AAC22387.1  (U32756) ATP-dependent DNA	1			
N=1~cDNA EST yk503h12.5 comes from this gene~cDNA EST yk653a11.3 comes from this gene~CDNA EST yk653a11.3 comes from this gene   The property   The property	1			
gene~cDNA EST yk653a11.3 comes from this gene   gene~cDNA EST yk653a11.5 comes from this gene   gene~cDNA EST yk653a11.5 comes from this gene   1.4	(			
gene~cDNA EST yk653a11.5 comes from this gene	ļ			
4880 7505750   Caenorha> 1.4				
gi[2500]12]sp[P71359]RECQ_HAEIN ATP- DEPENDENT DNA HELICASE RECQ gb]AAC22387.1] (U32756) ATP-dependent DNA	4880	7505750		1.4
DEPENDENT DNA HELICASE RECQ gb AAC22387.1  (U32756) ATP-dependent DNA				
gb AAC22387.1  (U32756) ATP-dependent DNA				
1002 2500112 picticase (1000) [riacinopinius influenzae Ku]	4882	2500112	helicase (recQ) [Hacmophilus influenzae Rd]	8.7

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 13447070 gb AAK26617.1  (AF340025) NADH	
4883	13447070	dehydrogenase subunit V [Coturnix chinensis]	3.1
		gi 14775307 ref XP_042250.1  similar to nuclear pore	
4886	14775307	complex interacting protein (H. sapiens) [Homo sapiens]	4.2
		gi 6434530 cmb CAB61080.1  (AL132949) predicted	
		using Genefinder~cDNA EST yk51g5.3 comes from this	
		gene~cDNA EST yk62d6.3 comes from this	
		gene~cDNA EST yk51g5.5 comes from this	
		gene~cDNA EST yk62d6.5 comes from this	
,		gene~cDNA EST yk105f8.5 comes from this	
4888	6434530	gene~cDNA EST yk99f8.3 >	0.28
		gi 12832739 dbj BAB22236.1  (AK002621) putative	
4896	12832739	[Mus musculus]	1.1
		gi 11281764 pir  C81739 conserved hypothetical protein	
		TC0120 [imported] - Chlamydia muridarum (strain	
4000	11201564	Nigg) gb AAF38998.1  (AE002279) conserved	
4903	11281764	hypothetical protein [Chlamydia muridarum]	2.7
4908	12020104	gi 12838194 dbj BAB24118.1  (AK005558) putative	
4908	12038194	[Mus musculus]	1.7
4911	12721121	gi 12721131 gb AAK02908.1  (AE006121) unknown [Pasteurella multocida]	0.2
4711	12/21131	gi 10638457 emb CAC12644.1  (AL136442) bA25J23.1	8.3
	*	gi 10638457 emb CAC12644.1  (AL136442) bA25J23.1 (KIAA1165 protein, similar to Drosophila CG8056	
4913	10638457	protein) [Homo sapiens]	1E-18
.,,13	10000437	gi 11346431 pir  T47803 mitogen-activated protein	112-10
1		kinase-like protein - Arabidopsis thaliana	
ĺ		emb CAB75798.1  (AL138647) mitogen-activated	
4916	11346431	protein kinase-like protein [Arabidopsis thaliana]	4
7710	115-0451	protein kinase-nae protein [Alabidopsis tilaitana]	

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundam	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gijl4758601 reffXP_010183.3  protein phosphatase, EF hand calcium-binding domain 1 [Homo sapiens] reffXP_040201.1  protein phosphatase, EF hand calcium-binding domain 1 [Homo sapiens] reffXP_040202.1  protein phosphatase, EF hand calcium-binding domain 1 [Homo sapiens] sp[014829 PEL_HUMAN SERINE/THREONINE PROTEIN PHOSPHATASE WITH EF-HANDS-1 (PPEF-1) (PROTEIN PHOSPHATASE WITH EF CALCIUM-BINDING DOMAIN) (PPEF) (SERINE/THREONINE PROTEIN PHOSPHATASE 7) (PP7) gb]AAB82795.1  (AF023455) protein phosphatase with EF-hands-1	
4924	14758601	[Homo sapiens] gb AAC05825.1  (AF027977) scrine/threonine protein phosphatase 7 catalytic subunit [Homo sapiens]	0.08
4932	12007365	gi 12007365 gb AAG45157.1 AF316823_1 (AF316823) cellulase Cel9-H [Clostridium cellulolyticum]	3.2
4933	8745051	gi 8745051 emb CAB95305.1  (AL359781) dynein heavy chain, cytosolic [Leishmania major]	8
4934	7499345	gil7499345 pir T21083 hypothetical protein F18A11.4 - Caenorhabditis elegans emb CAB04136.1  (Z81507) cDNA EST yk651h1.5 comes from this gene [Caenorhabditis elegans] emb CAA21639.1  (AL032639) cDNA EST yk651h1.5 comes from this gene [Caenorhabditis elegans]	0.4
4935	14133229	gi 14133229 dbj BAA76843.2  (AB023216) KIAA0999 protein [Homo sapiens]	0.91
4936		gij\$835447 reflNP_008371.1 ND6_13186 NADH dehydrogenase subunit 6 [Onchocerca volvulus] pil[711066 NADH dehydrogenase (ubiquinone) (EC 1.6.5.3) chain 6 - nematode (Onchocerca volvulus) mitochondrion gb[AAC61613.1] (AP015193) NADH dehydrogenase subunit 6 [Onchocerca volvulus]	2.8
4940		gij(6978439 ref]NP_036622.1  acrosin [Rattus norvegicus] emb CAA41441.2  (X58550) preproacrosin [Rattus sp.]	9.1

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 11349140 pir  G83492 hypothetical protein PA1232	
	[	[imported] - Pseudomonas aeruginosa (strain PAO1)	
		gb AAG04621.1 AE004552_11 (AE004552)	
4946	11349140	hypothetical protein [Pseudomonas aeruginosa]	1
		gi 12862434 dbj BAB32470.1  (AB047280) Pol-like	
4950	12862434	protein Pol-2 [Tricholoma matsutake]	4.6
		gi 15011489 gb AAK77584.1 AF396436_24	
4956	15011489	(AF396436) heme maturase [Tetrahymena thermophila]	2.8
		gi 1944163 dbj BAA19637.1  (AB002668) unnamed	
4958	1944163	protein product [Actinobacillus actinomycetemcomitans]	0.84
		W-50-100 / W-50-100	
		gi 7522108 pir  T29097 pro-pol-dUTPase polyprotein -	
1		murine endogenous retrovirus ERV-L (fragment)	
		emb CAA73251.1  (Y12713) protease; reverse transcriptase; RNaseH; integrase; dUTPase; Pro-Pol-	
4966	7522108	dUTPase; Rivaseri; integrase; dUTPase; Pro-Poi-	0.0005
4900	7322108	do i rase polyprotem [wius museums]	0.0003
		gi 12963743 ref NP_076085.1  equilibrative nucleoside	
1		transporter 3 [Mus musculus]	
		gb AAK00957.1 AF326986 1 (AF326986) equilibrative	
4968	12963743	nucleoside transporter 3 [Mus musculus]	9.3
		gi 6503300 gb AAF14676.1 AC011713 24 (AC011713)	
		Contains PF 01535 Domain of unknown function.	
4971	6503300	[Arabidopsis thaliana]	3.1
ļ.			
		gi 7498863 pir  T20730 hypothetical protein F10G8.8 -	
		Caenorhabditis elegans emb CAB02286.2  (Z80216)	
		Weak similarity with intermediate filament protein	
		(TREMBL id G633240), contains similarity to Pfam	
		domain: PF00169 (PH domain), Score=79.3, E-	
		value=2.7e-21, N=2~cDNA EST EMBL:T01262 comes	
		from this gene~cDNA EST yk23d5.3 comes from this	
		gene-> emb CAA19441.2  (AL023823) Weak similarity	
		with intermediate filament protein (TREMBL id	
		G633240), contains similarity to Pfam domain:	
		PF00169 (PH domain), Score=79.3, E-value=2.7e-21,	
1		N=2~cDNA EST EMBL:T01262 comes from this	
4976	7498863	gene~cDNA EST yk23d5.3 comes from this gen>	7

	l	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 436923 gb AAA62274.1  (U01849) ORF1	
4981	436923	[Trypanosoma brucei]	1.2
		gi 2827416 gb AAB99843.1  (AF043083) glycoprotein	
4982	2827416		9.1
		gi 11135234 sp P57642 SYK3_BUCAI PUTATIVE	
		LYSYL-TRNA SYNTHETASE (LYSINE-TRNA	
		LIGASE) (LYSRS) (GX) dbj BAB13271.1	
4000	11105004	(AP001119) hypothetical lysyl-tRNA synthetase	
4989	11135234	homolog [Buchnera sp. APS]	1.4
		gi 2129953 pir  JC5229 laccase (EC 1.10.3.2) precursor -	
4990	2120052	common tobacco gb AAC49536.1  (U43542) diphenol oxidase [Nicotiana tabacum]	9.1
4990	2129933	gi 13786443 gb AAK39568,1 AC025296 3	9.1
4995	13786443	(AC025296) hypothetical protein [Oryza sativa]	0.83
4993	13/80443	gi 7494262 pir T18489 hypothetical protein C0820w -	0.63
		malaria parasite (Plasmodium falciparum)	
	İ	emb CAB11128.1  (Z98551) hypothetical protein,	
4998	7494262	PFC0820w [Plasmodium falciparum]	0.81
		gi 10121788 gb AAG13373.1  (AF268180) polyprotein	
5000	10121788	[bovine viral diarrhea virus type 2]	7.4
		gi 6906704 dbj BAA90553.1  (AB032551) proline-rich	
		inositol polyphosphate 5-phosphatase [Rattus	
5007	6906704	norvegicus]	1.6
		gi 8978523 dbj BAA98360.1  (AP002545) CT147	
5008	8978523	hypothetical protein [Chlamydophila pneumoniae J138]	7.4
		gi 9630127 ref NP_046554.1  putative lipoprotein	
i		[Bacteriophage SPBc2] pir T12766 probable lipoprotein	
		yokB - Bacillus subtilis phage SPBc2 emb CAB14083.1	
		(Z99115) yokB [Bacillus subtilis] gb AAC12975.1	
5009	9630127	(AF020713) putative lipoprotein [Bacteriophage SPBc2]	2.7
3009	9030127	SF DC2]	
	l	gi 8978523 dbi BAA98360.1  (AP002545) CT147	
5011	8978523	hypothetical protein [Chlamydophila pncumoniae J138]	7.4
	-5.0525	gi 13636469 ref XP 016747.1  latent transforming	
		growth factor beta binding protein 1 precursor [Homo	
5012	13636469		1.7
		gi 3694664 gb AAC62434.1  (AC004893) similar to	
		NEDD-4 (KIA0093); similar to P46934	
5016	3694664	(PID:g1171682) [Homo sapiens]	7.9
		gi 14751816 ref XP_034239.1  hypothetical protein	
5020	14751816	XP_034239 [Homo sapiens]	0.064

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		"Marcon tol. 111 h Process all Process at the second	
5022	4768840	gi 4768840 gb AAD29638.1 AF117611_1 (AF117611)	
3022	4/68840	DosA protein [Dictyostelium discoidcum]	9.9
ĺ			
1		gi 7442182 pir  T04989 pathogenesis-related protein 1	
		precursor, 19.3K - Arabidopsis thaliana	
ł		emb CAA20585.1  (AL031394) pathogenesis-related	
		protein 1 precursor, 19.3K [Arabidopsis thaliana]	
		emb CAB80089.1  (AL161584) pathogenesis-related	
i		protein 1 precursor, 19.3K [Arabidopsis thaliana]	
]	J	gb AAG40056.1 AF324705_1 (AF324705) AT4g33720 [Arabidopsis thaliana] gb AAG42009.1 AF327419 1	
		[AF327419] putative pathogenesis-related protein 1	
		precursor protein [Arabidopsis thaliana]	
1		gb AAK00381.1 AF339699 1 (AF339699) putative	
	ł	pathogenesis-related protein 1 precursor protein	
		[Arabidopsis thaliana] gb AAK62632.1  (AY039577)	
5023	7442182	AT4g33720/T16L1 210 [Arabidopsis thaliana]	9.3
		gi 4884674 gb AAD31763.1 AF121945 1 (AF121945)	7.5
5029	4884674	aldehyde oxidase [Mus musculus]	7.9
		gi 2133751 pir  S66610 sulfakinin - bluebottle fly	
		(Calliphora vomitoria) prf  2120269A sulfakinin	
5031	2133751	[Calliphora vomitoria]	9
		gi 975667 emb CAA61500,1  (X89213) RNA	
		polymerase [Infectious hematopoictic necrosis virus]	
		prf  2121413F RNA polymerase [Infectious	
5033	975667	hematopoietic necrosis virus]	4.1
		gi 11267434 pir  F82853 DNA helicase II XF0050	
		[imported] - Xylella fastidiosa (strain 9a5c)	
		gb AAF82863.1 AE003859_4 (AE003859) DNA	
5035	11267434	helicase II [Xylella fastidiosa 9a5c]	3.1
		gi 11994373 dbj BAB02332,1  (AB019229)	
		gene_id:MDC16.14~unknown protein [Arabidopsis	
5036	11994373	thaliana]	4.8
		gi 1082778 pir  A56395 secretory phospholipase A2	
		receptor precursor, transmembrane form - human	
		gb AAA70110.1  (U17033) 180 kDa transmembrane	
5050	1082778	PLA2 receptor precursor [Homo sapiens]	3.5

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DECCRIPTION	B *** * * * * * * * * * * * * * * * * *
NO	IN .	DESCRIPTION	P VALUE
		gi 13518480 ref[NP 084839.1  hypothetical protein	
	ĺ	[Lotus japonicus] ref[NP 084856.1] hypothetical protein	
		[Lotus japonicus] dbj[BAB33238.1] (AP002983)	
		hypothetical protein [Lotus japonicus] dbj BAB33256.1	
5051	13518480	(AP002983) hypothetical protein [Lotus japonicus]	5
	10010100	gi 7460247 pir  B71612 hypothetical protein PFB0555c -	
		malaria parasite (Plasmodium falciparum)	
		gb AAC71900.1  (AE001402) hypothetical protein	
5065	7460247	[Plasmodium falciparum]	7.1
		gi 13486795 dbj BAB40028.1  (AP003021) hypothetical	
5069	13486795	protein [Oryza sativa]	2.6
		gi 6166049 sp O55017 CCAB MOUSE VOLTAGE-	
		DEPENDENT N-TYPE CALCIUM CHANNEL	
		ALPHA-1B SUBUNIT (CALCIUM CHANNEL, L	
		TYPE, ALPHA-1 POLYPEPTIDE ISOFORM 5)	
		(BRAIN CALCIUM CHANNEL III) (BIII)	
		gb AAB97840.1  (AF042317) neuronal type calcium	
5072	6166049	channel alpha-1 subunit [Mus musculus]	1.2
		gi 7495367 pir  T33395 hypothetical protein C04F2.1 -	
		Caenorhabditis elegans gb AAC26911.1  (AF078780)	
		Similar to chemoreceptor; C04F2.1 [Caenorhabditis	
5074	7495367	elegans]	3.1
		gi 9507739 ref NP_061405.1  23 pct identical(9 gaps) to	
		343 residues of 1286 aa protein	
		sp:AIDA_ECOLI[Adhesin AidA-I precursor] [Plasmid	
		F] dbj BAA97896.1  (AP001918) 23 pct identical(9	
		gaps) to 343 residues of 1286 aa protein	
5075	9507739	sp:AIDA_ECOLI[Adhesin AidA-I precursor] [Plasmid F]	0.7
3013	2301139	×1	9.7
		-1/7/1055091: 1/7190021	
		gi 7495508 pir  T18993 hypothetical protein C06B8.1 - Caenorhabditis elegans emb CAB03850.1  (Z81463)	
		Similarity to C.elegans zinc finger proteins, contains	
		similarity to Pfam domain: PF00104 (Ligand-binding	
	- 33	domain of nuclear hormone receptor), Score=-13.7, E-	
5076	7495508	value=0.051, N=1 [Caenorhabditis elegans]	4.2
		gi 14018037 ref NP 114377.1  ATP synthase F0 subunit	
		6 [Hymenolepis diminuta]	
		gb AAK51337,1 AF314223 9 (AF314223) ATP	
5077	14018037		4.9

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 6562005 emb CAB62499.1  (AJ250586) chitinase	
5086	6562005	[Arthrobacter sp.]	1.5
		gi 9631327 ref[NP_048161.1  ORF MSV090 putative	
		Molluscum contagiosum virus MC121L (vaccinia	
		A16L) homolog, similar to GB:U60315 [Melanoplus	
		sanguinipes entomopoxvirus] pir  T28251 ORF	
		MSV090 probable Molluscum contagiosum virus	
		MC121L (vaccinia A16L) homolog - Melanoplus	
		sanguinipes entomopoxvirus gb AAC97640.1	
		(AF063866) ORF MSV090 putative Molluscum	
		contagiosum virus MC121L (vaccinia A16L) homolog,	
		similar to GB:U60315 [Melanoplus sanguinipes	
5087	9631327	entomopoxvirus]	8.7
		gi 4321813 gb AAD15835.1  (AF063236) variant 2	
5088	4321813	major surface glycoprotein [Pneumocystis carinii]	6.5
5089	10050565	gi 12853765 dbj BAB29840.1  (AK015427) putative [Mus musculus]	
3069	12633763	[ivius musculus]	0.0003
		gi 732059 sp P39372 YJHA_ECOLI HYPOTHETICAL	
		28.3 KDA PROTEIN IN FECI-FIMB INTERGENIC	
		REGION PRECURSOR pir    S56536 hypothetical	
		protein f241 (fecI-fimB intergenic region) - Escherichia	
		coli gb AAA97207.1  (U14003) ORF_f241 [Escherichia	
		coli] gb AAC77267.1  (AE000501) orf, hypothetical	
		protein [Escherichia coli K12]	
		gb AAG59493.1 AE005662_5 (AE005662) orf,	
		hypothetical protein [Escherichia coli O157;H7 EDL933] dbj BAB38693.1  (AP002569) hypothetical	
5094	732059	protein [Escherichia coli O157:H7]	1.4
3034	152055	gi 84124 pir  B24785 hypothetical protein 1028 - slime	1,4
		mold (Dictyostelium discoideum) transposon DIRS-1	
5095	84124	(fragment)	2.9
3035	0.127	gi 84043 pir  C22845 NADH dehydrogenase	2.7
		(ubiquinone) (EC 1.6.5.3) chain 4 - Trypanosoma brucci	
5096	84043	mitochondrion	2.3
	2.315	gi 5441886 dbi BAA82384,1  (AP000367) EST	2.3
		AU069246(C53478) corresponds to a region of the	
5100	5441886	predicted gene.; hypothetical protein [Oryza sativa]	4.9

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
			1 11202
		gi 6678810 ref NP_032581.1  mannan-binding lectin	
	i	serine protease 1 [Mus musculus]	
		sp[P98064 CRAR_MOUSE COMPLEMENT- ACTIVATING COMPONENT OF RA-REACTIVE	
		FACTOR PRECURSOR (RA-REACTIVE FACTOR	
		SERINE PROTEASE P100) (RARF) (MANNAN-	
		BINDING LECTIN SERINE PROTEASE 1)	
		dbj BAA03944.1  (D16492) P100 serine protease of Ra-	
5102	6678810	reactive factor (RaRF) [Mus musculus]	7.3
		THOUSE COLUMN OF THE COLUMN OF	
		gi 13357651 ref[NP_077925.1  unique hypothetical membrane lipoprotein [Ureaplasma urealyticum]	
		pir  G82934 hypothetical protein UU094 [imported] -	
		Ureaplasma urealyticum gb AAF30500.1 AE002109 3	
		(AE002109) unique hypothetical membrane lipoprotein	
5106	13357651	[Ureaplasma urealyticum]	2.4
		gi 2499547 sp P77153 WZB_ECOLI PROBABLE	
		LOW MOLECULAR WEIGHT PROTEIN-	
		TYROSINE-PHOSPHATASE WZB pir  D64972	
		probable protein-tyrosine-phosphatase (EC 3.1,3,48)	
		wzb, low molecular weight - Escherichia coli	
		gb AAC77834.1  (U38473) putative acid phosphatase [Escherichia coli] gb AAC75122.1  (AE000296)	
		probable protein-tyrosine-phosphatase [Escherichia coli	
		K12] gb AAG57121.1 AE005432 2 (AE005432)	
		probable protein-tyrosine-phosphatase [Escherichia coli	
		O157:H7 EDL933] dbj BAB36289.1  (AP002560)	
		probable protein-tyrosine-phosphatase [Escherichia coli	
5108	2499547	O157:H7]	1.4
£100		gi 5292165 gb AAB01085.2  (U34402) single-subunit	
5109	5292165	RNA polymerase C [Triticum aestivum]	2.9
5112	2598890	gi 2598890 dbj BAA23297.1  (D86277) VP7 [Human rotavirus 3]	0.52
3112	2390090	gi 7301702 gb AAF56815.1  (AE003767) wdn gene	0.32
5115	7301702	product [Drosophila melanogaster]	2
		gi 12853765 dbj BAB29840.1  (AK015427) putative	
5116	12853765	[Mus musculus]	0.0003
		gi 4009428 gb AAD11553.1  (AF019894) replication	
5121	4009428	protein A [Helicobacter pylori]	4.2
5128	12940716	gi 12849716 dbj BAB28451.1  (AK012761) putative [Mus musculus]	7.3
3120	12049/10	[ivius musculus]	1.3

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS				
NO	N	DESCRIPTION	P VALUE		
		gi 7496182 pir  T19344 hypothetical protein C17D12.4 -			
1		Caenorhabditis elegans emb CAB03892.1  (Z81473)			
5129	7496182	predicted using Genefinder [Caenorhabditis elegans]	5.7		
		gi 9634540 ref NP_038078.1  TA35L [Vaccinia virus			
		(strain Tian Tan)] gb AAF34031.1  (AF095689) TA35L			
5130	9634540	[Vaccinia virus (strain Tian Tan)]	9.5		
		gi 12045115 ref[NP_072926.1  lipoprotein, putative			
		[Mycoplasma genitalium] sp[P47502 Y260_MYCGE			
1	ĺ	HYPOTHETICAL LIPOPROTEIN MG260			
		PRECURSOR pir  G64228 hypothetical protein			
		homolog MG260 - Mycoplasma genitalium			
1		gb AAC71481.1  (U39705) lipoprotein, putative			
5136	12045115	[Mycoplasma genitalium]	3.3		
		gi 2144796 pir  I36912 involucrin S - douroucouli			
		(fragment) gb AAA35376.1  (M25314) involucrin (small			
5138	2144796	allele) [Aotus trivirgatus]	4.3		
		gi 7509951 pir  T33900 hypothetical protein Y48A5A.1 -			
		Caenorhabditis elegans gb AAD12829.1  (AF125455)			
5140	7509951	Y48A5A.1 gene product [Caenorhabditis elegans]	1.3		
		gi 13701664 dbj BAB42957.1  (AP003135)			
		ORFID:SA1688~hypothetical protein, similar to			
		teichoic acid translocation ATP-binding protein tagH			
		[Staphylococcus aureus subsp. aureus N315]			
		dbj BAB58033.1  (AP003363) hypothetical protein			
5143	13701664	[Staphylococcus aureus subsp. aureus Mu50]	6.1		
		gi 7243223 dbj BAA92659.1  (AB037842) KIAA1421			
5147	7243223	protein [Homo sapiens]	6E-96		
		gi 14771907 ref XP_045594.1  hypothetical protein			
5148	14771907	MGC4816 [Homo sapiens]	5E-30		
		gi 6056374 gb AAF02838.1 AC009894_9 (AC009894)			
		Similar to serine/threonine kinases [Arabidopsis			
5163	6056374	thaliana]	9.5		

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 6322278 rcf NP_012352.1  mannosyltransferase	
		complex component; Mnn11p [Saccharomyces	
		cerevisiae] sp[P46985[YJS3_YEAST	
		HYPOTHETICAL 47.8 KD PROTEIN IN SWE1-	
		ATP12 INTERGENIC REGION pir    S56966 probable	
		membrane protein YJL183w - yeast (Saccharomyces	
		cerevisiae) emb CAA89478.1  (Z49458) ORF YJL183w	
5164	6322278	[Saccharomyces cerevisiae]	0.5
	l	gi 14738479 ref XP_027836.1  general transcription	
5168	14738479	factor IIIC, polypeptide 4 (90kD) [Homo sapiens]	77
		gi 4505321 ref NP_003862.1  myelin transcription factor	
		2; cerebrin-50 [Homo sapiens] pir  I52374 cerebrin-50 -	
	ļ	human gb AAB34231.1  (S76853) cerebrin-	
		50=cerebrospinal fluid protein [human, cerebral brain,	
5169	4505321	Peptide, 435 aa] [Homo sapiens]	5
	ĺ	gi 14336722 gb AAK61254.1 AE006464_22	
		(AE006464) Similar to pre-pro-megakarycyte	
5171	14336722	potentiating factor precursor [Homo sapiens]	0.22
		gi 13700778 dbj BAB42074.1  (AP003132)	
		ORFID:SA0834~hypothetical protein, similar to	
		lipopolysaccharide modification acyltransferase	
		[Staphylococcus aureus subsp. aureus N315]	
		dbj BAB57136.1  (AP003360) hypothetical protein	
5173	13700778	[Staphylococcus aureus subsp. aureus Mu50]	5.7
	l	gi 8894607 emb CAB94193.2  (AJ289710) envelope	
5174	8894607	protein [HERV-H/env60]	0.0002
		gi 14772400 ref XP_045944.1  hypothetical protein	
5176	14772400	XP_045944 [Homo sapiens]	2.5
		gi 8248741 gb AAB20211.2  (S61973) NMDA receptor	
5181	8248741	glutamate-binding subunit [Rattus sp.]	7.4
		gi 12329963 cmb CAC24680.1  (AL513062) possible	
5186	12329963	high molecular mass nuclear antigen [Leishmania major]	9.7
		"Igo150151 IO 45000 PRV4 Personner	
		gi 3915815 sp Q45032 PRIA_BORBU PRIMOSOMAL	
		PROTEIN N' (REPLICATION FACTOR Y)	
		pir  F70101 primosomal protein N (priA) homolog	
5187	3915815	Lyme disease spirochete gb AAC66393.1  (AE001115)	0.06
318/	2912012	primosomal protein N (priA) [Borrelia burgdorferi]	0.86

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 7516796 pir  H72575 hypothetical protein APE1888 -	
		Acropyrum pernix (strain K1) dbj BAA80893.1	
		(AP000062) 107aa long hypothetical protein	
5191	7516796	[Aeropyrum pernix]	9.9
		gi 4691710 gb AAD28038.1 AF119712_1 (AF119712)	
		bone morphogenetic protein BMP2/4 [Lytechinus	
5192	4691710	variegatus]	4.5
		gi 10639274 cmb CAC11276.1  (AL445063)	
		amylopullulanase related protein [Thermoplasma	
5198	10639274	acidophilum]	2.4
		gi 10998836 gb AAG26008.1 AF312017_1 (AF312017)	
5199	10998836	beta-glucosidase precursor [Tenebrio molitor]	3.1
		gi 7657228 rcf NP 055258.1  interleukin 17B [Homo	
		sapiens] ref[XP 003898.3  50367 [Homo sapiens]	
		refIXP 046987.1  interleukin 17B [Homo sapiens]	
		ref[XP 046984.1] interleukin 17B [Homo sapiens]	
		ref[XP 046985.1] interleukin 17B [Homo sapiens]	
		ref[XP 046986.1] 50372 [Homo sapiens]	
		gb AAF01318.1 AF184969 1 (AF184969) cytokine-like	
		protein ZCYTO7 [Homo sapiens]	
		gb AAF28104.1 AF152098 1 (AF152098) interleukin	
		17B [Homo sapiens] gblAAF78775.1 AF212311 1	
		(AF212311) interleukin 20 [Homo sapiens]	
		gb AAG39637.1 AF110385 1 (AF110385) interleukin-	
		17 beta; IL-17 beta [Homo sapiens]	
		gb AAG44136.1 AF218727 1 (AF218727) neuronal	
		interleukin-17 related factor [Homo sapiens]	
		gb AAK60336.1 AF386077_1 (AF386077) interleukin	
5205	7657228	17B [Homo sapiens]	5.5
		gi 12311878 emb CAC22694.1  (AL389894)	
5207	12311878	hypothetical protein L779.02 [Leishmania major]	1.6

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
NO	19	DESCRIPTION	P VALUE
	İ	gi 7657683 ref[NP_055146.1  solute carrier family 7,	
		(cationic amino acid transporter, y+ system) member 11;	
		cystine/glutamate transporter [Homo sapiens]	
Ì	}	sp Q9UPY5 XCT_HUMAN CYSTINE/GLUTAMATE	
		TRANSPORTER (AMINO ACID TRANSPORT	
		SYSTEM XC-) (XCT) dbj BAA82628.1  (AB026891)	
	1	cystine/glutamate transporter [Homo sapiens]	
		gb AAG35592.1 AF200708_1 (AF200708) calcium	
		channel blocker resistance protein CCBR1 [Homo	
		sapiens] gb AAK49111.1 AF252872_1 (AF252872)	
5208	7657683	cystine/glutamate transporter xCT [Homo sapiens]	2.2
		gi 2388574 gb AAB71455.1  (AC000098) Strong	
		similarity to Phalaenopsis homeobox protein	
5212	2388574	(gb U34743). [Arabidopsis thaliana]	7
		gi 14740481 ref XP_045136.1  similar to cadherin	
5221	14740481	related 23 (H. sapiens) [Homo sapiens]	1E-18
		gi 12852452 dbj BAB29417.1  (AK014534) putative	
5222	12852452	[Mus musculus]	8E-21
		gi 2119250 pir  I38857 microtubule-associated protein	
		1A - human gb AAA81362.1  (U14577) microtubule-	
5224	2119250	associated protein 1A [Homo sapiens]	0.99
		gi 1304610 gb AAC59915.1  (U41783) cytochrome b	
5226	1304610	[Cynolebias affinis]	4.7
_		gi 12311878 emb CAC22694.1  (AL389894)	
5227	12311878	hypothetical protein L779.02 [Leishmania major]	1.3
_		gi 6650047 gb AAF21692.1 AF051987_1 (AF051987)	
5228	6650047	maturase K [Hyobanche sanguinea]	1.9
		gi 3859671 emb CAA22009.1  (AL033502) hypothetical	
5232	3859671	protein [Candida albicans]	0.0009
		gi 9963891 gb AAG09748.1 AF233276_1 (AF233276)	
5240	9963891	peroxin-1 [Penicillium chrysogenum]	9.3
		gi 3024924 sp Q58315 Y905_METJA	
		HYPOTHETICAL PROTEIN MJ0905 pir  A64413	
		hypothetical protein MJ0905 - Methanococcus	
		jannaschii gb AAB98915.1  (U67534) M. jannaschii	
		predicted coding region MJ0905 [Methanococcus	
5242	3024924	jannaschii]	1.8
		oil11270021hid/T47006 hom shoti-ol acota i	
		gi 11279021 pir  T47996 hypothetical protein	
5245	11270021	F21F14.210 - Arabidopsis thaliana emb[CAB71911.1] (AL138642) putative protein [Arabidopsis thaliana]	
J243	112/9021	(AL130042) putative protein [Arabidopsis thaliana]	5.2

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 2746323 gb AAB97896.1  (AF037166) major surface	
5255	2746323	glycoprotein [Leishmania panamensis]	2.2
		gi 7506750 pir  T16770 hypothetical protein R153.2 -	
		Caenorhabditis elegans gb AAA68293.1  (U28729)	
5260	7506750	Hypothetical protein R153.2 [Caenorhabditis elegans]	4.1
		gi 14737112 ref XP_046972.1  hypothetical protein	
5264	14737112	XP_046972 [Homo sapiens]	3.4
		gi 7432465 pir  T13677 NADH dehydrogenase	
l i		(ubiquinone) (EC 1.6.5.3) chain 5 - Santolina	
1 1		chamaecyparissus chloroplast gblAAC37776.1	
		(L39444) NADH dehydrogenase [Santolina	
5266	7432465	chamaecyparissus]	8.1
		gi 9845030 dbj BAB11914.1  (AB034726) 5-	
5267	9845030	oxoprolinase precursor [Alcaligenes faecalis]	6.8
		gi 7429144 pir  GNMSLL retrovirus-related reverse	
		transcriptase homolog - mouse retrotransposon	
		gb AAA66024.1  (M13002) 2855 is the position of the	
5268	7429144	first start codon in ORF 2; putative [Mus musculus]	0.0003
		gi 6689319 emb CAB65444.1  (AJ238583) penicillin	0.0003
5269	6689319	binding protein 2x [Streptococcus pneumoniae]	0.37
		gi 7522099 pir  T28658 polyketide synthase - Sorangium	
5274	7522099	cellulosum (fragment)	3.9
		gi 13786182 ref NP 112634.1  delta-6 fatty acid	
		desaturase [Rattus norvegicus] pir JG0180 Delta6 fatty	
		acid desaturase (EC 1.14.99) [imported] - rat	
		dbi BAA75496.1  (AB021980) delta-6 fatty acid	
5278	13786182	desaturase [Rattus norvegicus]	4E-12
		gi 7674158 sp O76942 PTP ENCCU MAJOR POLAR	
		TUBE PROTEIN PRECURSOR (MAJOR PTP)	
		emb CAA06662.1  (AJ005666) polar tube protein	
5279	7674158	[Encephalitozoon cuniculi]	1.2
		gi 10726396 gb AAF54288.2  (AE003680) CG11773	
5280	10726396	gene product [Drosophila melanogaster]	1.1
		gi 14767779 ref XP_007023.2  G protein-coupled	
		receptor kinase-interactor 2 [Homo sapiens]	
		gb AAD28047.1 AF124491 1 (AF124491) ARF	
5282	14767779	GTPase-activating protein GIT2 [Homo sapiens]	4E-13
		gi 14602920 gb AAH09955.1 AAH09955 (BC009955)	
		Unknown (protein for IMAGE:4297851) [Homo	
5284	14602920		1.4
5284	14602920		1.4

	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS			
NO	N	DESCRIPTION	P VALUE	
		gi 7487876 pir  T02741 hypothetical protein T9I4.19 -		
	ļ	Arabidopsis thaliana gb AAC33237,1  (AC005315)		
		putative ligand-gated ion channel protein [Arabidopsis		
5287	7487876	thaliana]	9.9	
		gi 14423780 sp 095013 04F3_HUMAN OLFACTORY		
		RECEPTOR 4F3 gb AAD05195.1  (AC004908) similar		
		to rat olfactory receptor OR18; similar to S29710		
5288	14423780	(PID:g423702) [Homo sapiens]	0.000002	
		gi 14764967 ref XP_049605.1  KIAA1467 protein		
5290	14764967	[Homo sapiens]	0.00003	
1				
1		gi 7496712 pir  T15708 hypothetical protein C30B5.1 -		
ļ.		Caenorhabditis elegans gb AAK31466.1  (U23450)		
5291	7496712	Hypothetical protein C30B5.1 [Caenorhabditis elegans]	2.6	
		gi 4468835 emb CAB38221.1  (AJ232783) hairless		
5292	4468835	[Drosophila hydei]	1.3	
		gi 7481600 pir  T36589 probable transmembrane protein		
		- Streptomyces coelicolor emb CAB42730.1		
		(AL049826) putative transmembrane protein		
5298	7481600	[Streptomyces coelicolor A3(2)]	5.6	
J .		gi 4885323 ref NP 005293.1  G protein-coupled		
		receptor 37 (endothelin receptor type B-like); endothelin		
		receptor type B-like; hET(B)R-LP [Homo sapiens]		
		ref[XP_004804.1  G protein-coupled receptor 37		
1		(endothelin receptor type B-like) [Homo sapiens]		
		ref[XP_032150.1  58490 [Homo sapiens]		
		ref XP_032151.1  G protein-coupled receptor 37		
		(endothelin receptor type B-like) [Homo sapiens]		
		sp O15354 GP37_HUMAN PROBABLE G PROTEIN-		
1		COUPLED RECEPTOR GPR37 PRECURSOR		
		(ENDOTHELIN B RECEPTOR-LIKE PROTEIN-1)		
		(ETBR-LP-1) emb CAA73080.1  (Y12476) G protein		
		coupled receptor 37 [Homo sapiens] gb AAD08853.1		
		(AC004925) G protein coupled receptor 37 [Homo		
_5301	4885323	sapiens]	1.9	

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 11467524 ref[NP_043670.1  PSI, subunit III,	
		plastocyanin-binding [Odontella sinensis]	
		sp P49483 PSAF_ODOSI PHOTOSYSTEM I	
		REACTION CENTRE SUBUNIT III (PSI-F)	
		pir  S78329 photosystem I chain III - Odontella sinensis	
		chloroplast emb CAA91702.1  (Z67753) PSI, subunit	
5304	11467524	III, plastocyanin-binding [Odontella sinensis]	6.2
		gi 7292767 gb AAF48162.1  (AE003489) CG15927	
5306	7292767	gene product [Drosophila melanogaster]	6.3
		gi 14601766 ref NP_148307.1  hypothetical protein	
		[Aeropyrum pernix] pir  C72501 hypothetical protein	
		APE1985 - Aeropyrum pernix (strain K1)	
		dbj BAA80995.1  (AP000063) 160aa long hypothetical	
5309	14601766	protein [Aeropyrum pernix]	1.8
		gi 13661965 gb AAK38127.1 AC058781_4	
5310	13661965	(AC058781) L344.4 [Leishmania major]	7.1
		gi 6320879 ref NP_010958.1  Transcriptional activator	
		of nitrogen-regulated genes; Gln3p [Saccharomyces	
		cerevisiae] sp P18494 GLN3_YEAST NITROGEN	
		REGULATORY PROTEIN GLN3 pir S50543 GLN3	
		protein - yeast (Saccharomyces cerevisiae)	
5010	6320879	gb AAB64575.1  (U18796) Gln3p: Nitrogen regulatory	0.5
5312	6320879	protein [Saccharomyces cerevisiae]	8.5
		gi 4493990 emb CAB39049.1  (AL034559) hypothetical	
5313	4493990	protein, PFC1045c [Plasmodium falciparum]	0.74
3313	4493990	gi 10047245 dbi BAB13411.1  (AB046805) KIAA1585	0.74
5314	10047245	protein [Homo sapiens]	2E-69
3314	10047243	gi 14762995 ref XP 044123.1  cadherin 20, type 2	2E-09
5317	14762995		4E-17
3317	14702333	[riomo sapiens]	4E-17
5321	14773348	gi 14773348 ref XP_038450.1  20849 [Homo sapiens]	3E-45
		gi 4691710 gb AAD28038.1 AF119712 1 (AF119712)	02.0
	1	bone morphogenetic protein BMP2/4 [Lytechinus	
5326	4691710	variegatus	5.4
		gi 13161382 dbj BAB32977,1  (AB034197) lamin B3	
5328	13161382		6.9
		gi 12654811 gb AAH01248.1 AAH01248 (BC001248)	
5333	12654811	hypothetical protein FLJ20272 [Homo sapiens]	6.2

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	· DESCRIPTION	P VALUE
		gi 4504455 ref NP_000183.1  T-box 5 [Homo sapiens]	
l		emb CAA70592.1  (Y09445) transcription factor [Homo	
5334	4504455	sapiens]	2.5
		  gi 6642649 gb AAF20230.1 AC012395 17 (AC012395)	
5337	6642649	putative glucan synthase [Arabidopsis thaliana]	1.7
- 3337	0042049	putative giucan symmase [Arabidopsis manana]	1.7
		gi 7330072 gb AAF60061.1 AF210726 82 (AF210726)	
5342	7330072		0.012
		gi 90587 pir  PS0135 H-2 class I histocompatibility	
		antigen T7 - mouse (fragment) emb CAA34332.1	
		(X16213) MHC T7 class I antigen (64 AA) (119 is 2nd	
5344	90587	base in codon) [Mus musculus]	6.4
		gi 14749813 ref XP_041197.1  integrin, alpha 11 [Homo	
5345	14749813	sapiens]	0.48
		gi 2143962 pir  159422 rsec8 - rat (fragment)	
5348	2143962	gb AAC52265.1  (U32498) rsec8 [Rattus norvegicus]	3E-96
		gi 7549797 ref NP_035731.1  T lymphoma oncogene	
		[Mus musculus] sp P17408 TLM_MOUSE TLM	
		PROTEIN (TLM ONCOGENE) pir   S10151	
	*	transforming protein tlm - mouse (strain balb/c)	
5349	7549797	emb CAA36859.1  (X52634) tlm protein [Mus musculus]	1.9
3349	1343131	inusculusj	1.9
		gi 7505043 pir  T33641 hypothetical protein K01A2.7 -	
		Caenorhabditis elegans gb AAC69507.1  (AF099925)	
5351	7505043	Hypothetical protein K01A2.7 [Caenorhabditis elegans]	0.71
		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	
		gi 6677735 ref NP 033084.1  ral guanine nucleotide	
		dissociation stimulator [Mus musculus]	
		sp Q03385 GNDS MOUSE RAL GUANINE	
		NUCLEOTIDE DISSOCIATION STIMULATOR	
		(RALGEF) (RALGDS) pir   S28415 guanine nucleotide	
		dissociation stimulator ralGDS - mouse	
l		gb AAA37714.1  (L07924) guanine nucleotide	
5357	6677735	dissociation stimulator [Mus musculus]	4
		gi 11499595 ref[NP_070837.1  coenzyme F390	
		synthetase (ftsA-3) [Archaeoglobus fulgidus]	
		pir  D69501 coenzyme F390 synthetase (ftsA-3)	
		homolog - Archaeoglobus fulgidus gb AAB89243.1  (AE000964) coenzyme F390 synthetase (ftsA-3)	
5360	11499595	[Archaeoglobus fulgidus]	4.8
2200	***//////	[1 TOWNOOPIODON TRIBUTAD]	

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS		110,0000)	
NO	N	DESCRIPTION	P VALUE	
5365	12383066	gi[12383066]ref[NP_073737.1] hypothetical protein DKFZp586F1122 similar to axotrophin [Homo sapiens] ref[XP_016117.2] hypothetical protein DKFZp586F1122 similar to axotrophin [Homo sapiens] dbj[BAB14340.1] (AK022973) unnamed protein product [Homo sapiens]	8E-61	
		gi 7500833 pir T21991 hypothetical protein F39B2.10 - Caemorhabditis elegans emb CAB07390.1  (292834) contains similarity to Pfam domain: PF00226 (DnaJ domain), Score=132.3, E-value=2.8e-36, N=1; PF00684 (DnaJ central domain (4 repeats)), Score=103.9, E-value=9.9e-28, N=1; PF01556 (DnaJ C terminal region), Score=35.8, E-value=1.5e-08,		
5368	7500833	N=1-cDNA> gi 11499595 ref[NP_070837.1  coenzyme F390 synthetase (ftsA-3) [Archaeoglobus fulgidus] pir[D69501 coenzyme F390 synthetase (ftsA-3) homolog - Archaeoglobus fulgidus gb AAB89243.1  (AE000964) coenzyme F390 synthetase (ftsA-3)	3.2	
5370	11499595		5.5	
5371	11387290	gi 11387290 sp P57436 Y355_BUCAI PUTATIVE DEOXYRIBONUCLEASE BU355 dbj BAB13059.1  (AP001119) hypothetical protein [Buchnera sp. APS]	7.1	
5374	8923462	gi 8923462 ref[NP_060317.1  hypothetical protein FLJ20505 [Homo sapiens] dbj BAA91218.1  (AK000512) unnamed protein product [Homo sapiens]	5E-76	
5376	7491381	gi 7491381 pir  T39498 hypothetical protein SPBC1604.16c - fission yeast (Schizosaccharomyces pombe) emb CAA22349.1  (AL034433) hypothetical protein [Schizosaccharomyces pombe]	5.1	
5378	5835229	gij5835229[ref]NP_008273.1]ND4_10703 NADH dehydrogenase subunit 4 [Protopterus dolloi] pirj568137 NADH dehydrogenase (ubiquinore) (EC 1.6.5.3) chain 4 - Protopterus dolloi mitochondrion gb[AAC38030.1] (L42813) NADH dehydrogenase subunit 4 [Protopterus dolloi]	7,9	
5381		gi 12861366 dbj BAB32182.1  (AK020701) putative [Mus musculus]	1E-20	

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ ID NO	ACCESS N	DESCRIPTION	DAVALTE		
NO	11	gi 12957488 ref NP_075463.1  putative gene product	P VALUE		
		[Homo sapiens] emb CAB42442.1  (AL049784)			
5387	12957488	hypothetical protein [Homo sapiens]	1E-23		
		-yp	10 25		
		gi 13621158 ref NP 112433,1  NADH dehydrogenase			
		subunit 1 [Tetrodontophora bielanensis]			
		gb AAK30952.1 AF272824 13 (AF272824) NADH			
5390	13621158	dehydrogenase subunit 1 [Tetrodontophora bielanensis]	8.5		
		gi 14720363 ref XP_042844.1  similar to ALU			
5201	1.45000.00	SUBFAMILY SC SEQUENCE CONTAMINATION			
. 5391	14720363	WARNING ENTRY (H. sapiens) [Homo sapiens]	5.5		
		gi 14916308 gb AAK73874.1 U00067 3 (U00067)			
5399	14916308	Hypothetical protein F54E7.3a [Caenorhabditis elegans]	7.6		
1	11,710000	gi 12856615 dbi BAB30727.1  (AK017396) putative	7.0		
5401	12856615		1E-91		
		gi 625580 pir  A49626 transregulatory protein IE-1 -			
		Autographa californica nuclear polyhedrosis virus			
		gb AAB29676.1  (S68091) IE-1=transregulatory protein			
1		[Autographa californica nuclear polyhedrosis virus			
5406	(25500	AcNPV, tsB821, Peptide Mutant, 582 aa] [Autographa californica nucleopolyhedrovirus]	20		
3406	625580	californica nucleopolyhedrovirus]	2.9		
		gi 7511058 pir  T27805 hypothetical protein ZK262,11 -			
		Caenorhabditis elegans emblCAB16552.1  (Z99288)			
		contains similarity to Pfam domain: PF01604 (7TM			
i i		chemoreceptor), Score=138.5, E-value=3.8e-38, N=1			
5408	7511058	[Caenorhabditis elegans]	8		
		gi 5869818 cmb CAB55575.1  (AJ249395) NADH-			
		ubiquinone oxidoreductase subunit 6 [Globodera			
5412	5869818	pallida]	0.36		
		gi 7492086 pir  T41670 hypothetical zinc finger protein -			
		fission yeast (Schizosaccharomyces pombe) emb CAA20703.1  (AL031530) hypothetical zinc finger			
<u> -  </u>		protein; C3HC4 type (RING finger) family			
5413	7492086	Schizosaccharomyces pombel	8.5		
	5000	gi 6056374 gb AAF02838.1 AC009894 9 (AC009894)			
		Similar to serine/threonine kinases [Arabidopsis			
5416	6056374	thaliana]	9.5		

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 7464490 pir  E64666 hypothetical protein HP1173 -	
		Helicobacter pylori (strain 26695) gb AAD08225.1	
		(AE000623) H. pylori predicted coding region HP1173	
5417	7464490	[Helicobacter pylori 26695]	4
		gi 13812393 ref NP 113511.1  hypothetical protein	
		[Guillardia theta] cmb CAC27080.1  (AJ010592)	
5418	13812393	hypothetical protein [Guillardia theta]	1.9
		· · · · · · · · · · · · · · · · · · ·	
		gi 6041669 ref NP_004538.2  NADH dehydrogenase	
	i	(ubiquinone) 1 beta subcomplex, 4 (15kD, B15) [Homo	
1	ļ	sapiens] ref[XP_002929.1  NADH dehydrogenase	
	1	(ubiquinone) 1 beta subcomplex, 4 (15kD, B15) [Homo	
i l		sapiens] ref[XP_041367.1  NADH dehydrogenase	
,		(ubiquinone) 1 beta subcomplex, 4 (15kD, B15) [Homo	
1	1	sapiens] sp O95168 NB5M_HUMAN NADH-	
		UBIQUINONE OXIDOREDUCTASE B15 SUBUNIT	
	ļ	(COMPLEX I-B15) (CI-B15) pir  JE0383 NADH	
		dehydrogenase (ubiquinone) (EC 1.6.5.3) chain	
		NDUFB4 - human gb AAD05421.1  (AF044957)	
1		NADH:ubiquinone oxidoreductase B15 subunit [Homo	
1	l	sapiens] gb AAH00855,1 AAH00855 (BC000855)	
		NADH dehydrogenase (ubiquinone) 1 beta subcomplex,	
5420	6041669	4 (15kD, B15) [Homo sapiens]	0.00002
		gi 7296624 gb AAF51905.1  (AE003600) CG10303	
5422	7296624	gene product [Drosophila melanogaster]	4.2
		gi 6453299 emb CAA04499.2  (AJ001045) P-type	
5429	6453299	cation-transporting ATPase [Blastocladiella emersonii]	6.2
		gi 14784393 ref XP_033306.1  sodium channel,	
5430	14784393	nonvoltage-gated 1 alpha [Homo sapiens]	7.6
		gi 7657956 dbj BAA94876.1  (AB028668) ORF1 [TT	
5438	7657956		3.4
	1	gi 7206840 gb AAF39999.1  (AC006832) similar to a	
		family of C. elegans proteins; see (GB:AF016684)	
5449	7206840	[Caenorhabditis elegans]	6.3
		gi 13636619 ref XP_002437.3  cAMP-regulated guanine	
5451	13636619	nucleotide exchange factor II [Homo sapiens]	0.000001
1		gi 7303178 gb AAF58242.1  (AE003814) CG17390	
5454	7303178	gene product [Drosophila melanogaster]	3.1

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Destains)
SEO ID	ACCESS	Table 36 Nearest Neighbor (Blasta vs. Non-Redundant	Proteins)
NO	N	DESCRIPTION	P VALUE
NO	IN		P VALUE
		gi 7446747 pir  T07607 phosphate transport protein 2	
5457	7446747	potato emb CAA67396.1  (X98891) inorganic phosphate transporter 2 [Solanum tuberosum]	4.0
3437	/446/4/	transporter 2 [Solanum tuberosum]	4.2
		-174712061-111475450	
		gi 7471296 pir  A75450 conserved hypothetical protein - Deinococcus radiodurans (strain R1)	
		gb AAF10584.1 AE001952_12 (AE001952) conserved	
5458	7471206	hypothetical protein [Deinococcus radiodurans]	0.32
3436	7471290	hypotheticai protein [Deinococcus radiodurans]	0.32
		"110/71501 : #401000 FF 77 1 1 1 4 mp	
		gi 11267150 pir  A81338 H+/K+-exchanging ATPase	
		(EC 3.6.1.36) B chain Cj0677 [imported] - Campylobacter jejuni (strain NCTC 11168)	
		emb CAB72951.1  (AL139076) potassium-transporting	
5460	11267150	ATPase B chain [Campylobacter jejuni]	0.37
3400	1120/130	gi 1246530 emb CAA64368,1  (X94742) olfactory	0.37
5467	1246530	receptor 2 [Gallus gallus]	0.00000001
3407	1240330	gil1353257 gb AAB06234,1  (U26665) dimethyl	0.0000001
		sulphoxide reductase subunit B [Haemophilus	
5475	1353257	influenzael	4.2
1 3473	1333237	initationizatej	7.2
		"Second Inageneining Course & STROME	
		gi 585053 sp P37202 DIS3_SCHPO MITOTIC CONTROL PROTEIN DIS3 pir  A41944 mitotic	
		control protein dis3+ - fission yeast	
		(Schizosaccharomyces pombe) gb AAA35302.1	
		(M74094) mitotic control protein [Schizosaccharomyces	
ļ		pombe] emb[CAA21102.1] (AL031743) mitotic control	
5476	585053	protein dis3 [Schizosaccharomyces pombe]	2.6
3470	363033	protein disc [Scinzosacciaroniyees pointe]	2.0
i			
		-372992191- JOOGLOTETHE LIEUDI DEODADI E	
		gi 7388318 sp Q9ZL01 THIE_HELPJ PROBABLE THIAMINE-PHOSPHATE PYROPHOSPHORYLASE	
		(TMP PYROPHOSPHORYLASE) (TMP-PPASE)	
		(THIAMINE-PHOSPHATE SYNTHASE) pir  G71889	
		thiamin-phosphate pyrophosphorylase (EC 2.5.1.3) -	
		Helicobacter pylori (strain J99) gb AAD06361.1	*
ĺ		(AE001508) THIAMINE PHOSPHATE	
5477	7388318	PYROPHOSPHORYLASE [Helicobacter pylori J99]	7
	.500510	pylotisss [noncomma pylotisss]	<del></del>
l		gi 6754362 ref NP 035962.1  insulin receptor-related	
		receptor [Mus musculus] dbi BAA77835.1  (AB007135)	
5483	6754362	insulin receptor-related receptor [Mus musculus]	7.5
5.05	1 1502	mount receptor remove receptor [mas museums]	

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	3 \	
NO	N	DESCRIPTION	P VALUE
		gi 10697032 emb CAC12661.1  (AJ296086) cytochrome	
5486	10697032	c oxidase subunit 1 [Anabaena variabilis]	0.96
1		gi 8919908 emb CAB96228.1  (AJ133067) cnv	
5491	8919908	polypeptide [Human immunodeficiency virus type 1]	7.4
	[	gi 7658073 dbj BAA94924.1  (AB028715) ORF1 [TT	
5496	7658073	virus]	3.8
		gi 9437326 gb AAF87312.1 AF124441_1 (AF124441)	
5500	9437326	NBC-like protein [Rattus norvegicus]	6.5
		gi 6707844 gb AAF25692.1  (AF113517) ventral	
5501	6707844	anterior homeobox 3 [Xenopus laevis]	3.1
		gi 12698075 dbj BAB21856.1  (AB051552) KIAA1765	
5502	12698075	protein [Homo sapiens]	1.9
		gi 7673065 gb AAF66723.1 AF146723_1 (AF146723)	
5503	7673065	cytochrome b [Tapinoma sp. ACBJ.1]	4.2
		gi 6754932 ref NP_035121.1  olfactory receptor 49	
5500	C754022	[Mus musculus] gb AAD13315.1  (AF102523) olfactory	07.10
5508	6/54932	receptor C6 [Mus musculus] gil7296803[gb AAF52080.1] (AE003604) CG2008 gene	8E-12
5512	7296803	gij/296803[gb]AAF52080.1] (AE003604) CG2008 gene product [Drosophila melanogaster]	3.7
3312	7290603		3.1
Ì		gi 5824783 cmb CAB54433.1  (AL110487) contains similarity to Pfam domain: PF01529 (DHHC zinc finger	
	}	domain), Score=108.2, E-value=5e-29, N=1~cDNA	
		EST vk282d12.5 comes from this gene [Caenorhabditis	
5514	5824783	elegansl	4.9
3314	3024703	cicgansj	4.7
1	ĺ	  gi 14750790 ref XP_034677.1  general transcription	
		factor II, i, isoform 5 [Homo sapiens] ref[XP 011605.4]	
		general transcription factor II. i. isoform 5 IHomo	
		sapiens] ref[XP 034678.1] similar to general	
ļ		transcription factor II, i (H. sapiens) [Homo sapiens]	
		ref[XP_034682.1  similar to general transcription factor	
5516	14750790	II, i (H. sapiens) [Homo sapiens]	4.2
		,, ,, ,, ,, ,, ,, ,, ,, ,, ,,	
		gi 3327362 dbi BAA31704.1  (AB015754) cytochrome c	
5519	3327362	oxidase subunit I [Spirometra erinaccieuropaei]	5.2
		gi 14149807 ref NP_115517.1  hypothetical protein	
		DKFZp434K1421 [Homo sapiens] emb CAB66740.1	
5520	14149807	(AL136806) hypothetical protein [Homo sapiens]	5E-53

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESS	The state of the s	1 Totoliis)		
NO	N	DESCRIPTION	P VALUE		
		gi 10567777 gb AAG18583.1 AF292395 1 (AF292395)			
5521	10567777	KRP170 [Strongylocentrotus purpuratus]	8.2		
		gi 5931705 emb CAB56603.1  (Y18890) pol protein			
5524	5931705	[Human endogenous retrovirus K]	6.2		
		gi 11271464 pir  C82792 methionyl-tRNA synthetase			
		XF0549 [imported] - Xylella fastidiosa (strain 9a5c)			
		gb AAF83359.1 AE003902_4 (AE003902) methionyl-			
5528	11271464	tRNA synthetase [Xylella fastidiosa 9a5c]	6.5		
		gi 2193879 cmb CAA54180.1  (X76785) hypothetical			
5530	2193879	protein [Homo sapiens]	8.7		
		gi 7292567 gb AAF47967.1  (AE003484) CG2186 gene	4.		
5531	7292567	product [Drosophila melanogaster]	6.7		
l	ŀ	gi 729093 sp P39881 CUT1_CANFA CCAAT			
		DISPLACEMENT PROTEIN (HOMEOBOX			
		PROTEIN CLOX) (CLOX-1) pir  S33121 homeotic			
5532	729093	protein CDP - dog (fragment) emb CAA48782.1  (X69017) Clox [Canis sp.]	0.14		
3332	129093	gi 13569850 ref NP 076357.1  RIKEN cDNA	0.14		
		9330127I20 [Mus musculus] gb AAG34081.1			
		(AF295105) cardiac Ca2+ release channel [Mus			
5534	13569850	musculus]	0.00000006		
	15503050	maseurasj	0.0000000		
		gi 14250321 gb AAH08590.1 AAH08590 (BC008590)			
5535	14250321	hypothetical protein FLJ21313 [Homo sapiens]	1E-60		
		gi 2143962 pir  I59422 rsec8 - rat (fragment)			
5536	2143962	gb AAC52265.1  (U32498) rsec8 [Rattus norvegicus]	3E-99		
		gi 13475196 ref NP_106760.1  transposase			
1	ł	[Mesorhizobium loti] dbj BAB52546.1  (AP003008)			
5539	13475196	transposase [Mesorhizobium loti]	5.6		
		gi 13376066 ref NP_079019.1  hypothetical protein			
		FLJ21934 [Homo sapiens] ref[XP_017673.2]			
	ļ	hypothetical protein FLJ21934 [Homo sapiens]			
		dbj BAB15179.1  (AK025587) unnamed protein product			
5544	13376066	[Homo sapiens]	1E-40		
		gi 13365831 dbj BAB39301.1  (AB056753) hypothetical			
5545	13363831	protein [Macaca fascicularis] gil2865238lgblAAC38862.1l (U89706) DNA	4.7		
5547	2865238	gi[2865238[gb]AAC38862.1] (U89706) DNA polymerase alpha [Urostyla grandis]			
3347	2003238	gi 14752983 reffXP 004626.3  hypothetical protein	5.5		
5548	14752993	FLJ10377 [Homo sapiens]	2.5		
3346	14132303	LEGIOSTI [LIOHO Sapicis]	2.3		

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 13489060 ref NP_109590.1  retinoic acid induced 1;	
		hypothetical protein DKFZp434A139 [Homo sapiens]	
		emb CAC20423.1  (AJ271790) retinoid-acid induced	
5550	13489060	protein 1 [Homo sapiens]	1E-21
		gi 7512077 pir  T30878 dyncin heavy chain isotype 4 -	
		sea urchin (Tripneustes gratilla) (fragment)	
		gb AAA63587.1  (U03973) dynein heavy chain isotype	
5556	7512077	4 [Tripneustes gratilla]	4E-27
		gi 9971630 dbj BAB12582.1  (AB046525) polymerase	
5564	9971630	protein [Hepatitis B virus]	9.1
		gi 1072920 pir  S49369 mobilization protein -	
		Campylobacter coli plasmid pCCT1 and PCCT2	
		emb CAA57597.1  (X82080) Mob [Campylobacter coli]	
		emb CAA57594.1  (X82079) mobilization protein	
5565	1072920	[Campylobacter coli]	4.7
		gi 12644030 sp Q28295 VWF_CANFA VON	
5568	12644030	WILLEBRAND FACTOR PRECURSOR (VWF)	8.8
		gi 9964333 ref NP_064801.1  AMV019 [Amsacta	
		moorei entomopoxvirus] gb AAG02725.1 AF250284_19	
		(AF250284) AMV019 [Amsacta moorei	
5572	9964333	entomopoxvirus]	8.3
		gi 10436768 dbj BAB14906.1  (AK024391) unnamed	
5579	10436768	protein product [Homo sapiens]	1.2
		gi 13786443 gb AAK39568.1 AC025296_3	
5580	13786443	(AC025296) hypothetical protein [Oryza sativa]	0.75
		gi 13376632 ref NP 079355.1  hypothetical protein	
		FLJ23231 [Homo sapiens] dbj[BAB15581.1]	
		(AK026884) unnamed protein product [Homo sapiens]	
		gb AAH05001.1 AAH05001 (BC005001) hypothetical	
5581	13376632	protein FLJ23231 [Homo sapiens]	0.58
		gi 12641960 gb AAK00073.1 AF200382 1 (AF200382)	
5584	12641960	cytochrome oxidase subunit I (COI) [Ceratosolen nanus]	4.9
		gil 1708893 splP51782 LYC TRIVU LYSOZYME C	
		PRECURSOR (1,4-BETA-N-	
		ACETYLMURAMIDASE C) gb AAB97109.1	
5585	1708893	(U40664) lysozyme [Trichosurus vulpecula]	2.3
		gi 10047155 dbj BAB13371.1  (AB046765) KIAA1545	
5586	10047155	protein [Homo sapiens]	1.7

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS			
NO	N	DESCRIPTION	P VALUE	
		gi 7446806 pir  F69791 conserved hypothetical protein		
l .		yebB - Bacillus subtilis gb AAB62312.1  (U51115)		
		unknown protein [Bacillus subtilis] emb CAB12456.1		
		(Z99107) similar to hypothetical proteins [Bacillus		
5587	7446806	subtilis]	9.3	
l				
		gi 400222 sp P22533 MANB CALSA BETA-		
		MANNANASE/ENDOGLUCANASE A		
		PRECURSOR [INCLUDES: MANNAN ENDO-1,4-		
	-	BETA-MANNOSIDASE A (BETA-MANNANASE)		
		(ENDO-1,4-MANNANASE); ENDO-1,4-BETA-		
		GLUCANASE (CELLULASE)] pir  A48954 mannan		
		endo-1,4-beta-mannosidase (EC 3.2.1.78) - Caldocellum		
		saccharolyticum gb AAA71887.1  (L01257) beta-		
5591	400222	mannanase [Caldicellulosiruptor saccharolyticus]	8.6	
		gi 14029388 gb AAK52669.1 AF319948_1 (AF319948)		
5593	14029388	MMS19 [Drosophila melanogaster]	7.7	
		gi 1196482 gb AAA88209.1  (M20307) unknown		
5595	1196482	protein [Plasmid P1]	5.2	
		gi 14762995 ref XP_044123.1  cadherin 20, type 2		
5599	14762995	[Homo sapiens]	4E-17	
		gi 13383732 gb AAK21107.1 AF327877_3 (AF327877)		
5600	13383732	envelope polyprotein [Equine infectious anemia virus]	3.7	
		gi 7521539 pir  A70410 processing proteinase (EC 3.4		
		) - Aquifex aeolicus gb AAC07272.1  (AE000732)		
5611	7521539	processing protease [Aquifex aeolicus]	9.5	
5612	14721452	-111472145210VD 040512 1148204 FT	,	
5612	14/21452	gi 14721452 ref XP_049513.1  48294 [Homo sapiens] gi 13512594 gb AAK28688.1  (AF078553) unknown	1	
5615	12512504	function U3 [Ehrlichia canis]	9.1	
3013	13512594		9.1	
		gi 7490184 pir  T37997 carboxypeptidase y - fission		
1		yeast (Schizosaccharomyces pombe) pir T43236		
		carboxypeptidase C (EC 3.4.16.5) precursor [validated]		
1		fission yeast (Schizosaccharomyces pombe)		
1	*	emb CAB10121.1  (Z97209) carboxypeptidase y		
		[Schizosaccharomyces pombe] dbj BAA25568.1		
5618	7490184	(D86560) carboxypeptidase Y [Schizosaccharomyces	0.01	
3010	7490184	gil1119139 gb AAG30518.1  (AF308549)	0.01	
5619	11110120	immunoglobulin heavy chain [Homo sapiens]	1.1	
2019	11119139	minimogroudin neavy cham [rionio sapiens]	1.1	

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	D. A.S. A
SEQ ID	ACCESS	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
NO NO	N N	DESCRIPTION	P VALUE
110	- 13	DESCRIPTION	FVALUE
		gi 7507209 pir  T29623 hypothetical protein T05E8.3 -	
		Caenorhabditis elegans gb[AAB52427,1] (U97014)	
	İ	strong similarity to the 'DEAH' subfamily of the 'DEAD'	
5622	7507209	box family of helicases [Cacnorhabditis elegans]	0.7
JULL	7307209	gi 7484413 pir  T08026 hypothetical protein B -	0.7
		Chlamydomonas reinhardtii chloroplast	
		gb AAB05800.1  (U62943) unknown [Chlamydomonas	
5623	7484413	reinhardtii]	9,3
3023	7404413	gi 14766128 ref XP 038109.1  KIAA1683 protein	7.3
		[Homo sapiens] emb CAB66801.1  (AL136867)	
5624	14766128	hypothetical protein [Homo sapiens]	0.15
	11700120	gi 6491915 gb AAF14073,1 AF104231 1 (AF104231)	0.13
		paired-homeodomain transcription factor PAX4 [Mus	
5629	6491915	musculus]	2.8
	0.077710		2.0
		gi 14149940 ref NP 115610.1  hypothetical protein	
		FLJ23059 [Homo sapiens] dbj BAB15536.1	
5630	14149940	(AK026712) unnamed protein product [Homo sapiens]	1E-10
		gi 7500466 pir  T21747 hypothetical protein F35C12.2 -	
5636	7500466	Caenorhabditis elegans	3.7
		gi 7490377 pir  T41496 conserved hypothetical protein	·
		SPCC622.16c - fission yeast (Schizosaccharomyces	
		pombe) emb CAA21872.1  (AL033127) conserved	
5638	7490377	hypothetical protein [Schizosaccharomyces pombe]	6.3
		gi 6322907 ref NP 012980.1  heavy chain of	
		cytoplasmic dynein; Dyn1p [Saccharomyces cerevisiae]	
		sp[P36022]DYHC_YEAST DYNEIN HEAVY CHAIN,	
		CYTOSOLIC (DYHC) pir S38128 dynein heavy chain,	
		cytosolic - yeast (Saccharomyces cerevisiae)	
		emb CAA82132.1  (Z28279) ORF YKR054c	
5642	6322907	[Saccharomyces cerevisiae]	4
		gi 7594617 cmb CAB88111.1  (AL078581) dJ12G14.1	
		(novel cyclophilin type peptidyl-prolyl cis-trans	
5646	7594617	isomerase) [Homo sapiens]	8E-73
		gi 14916439 ref NP_149094.1  rhophilin-like protein	
	l	[Homo sapiens] gb AAK58588.1 AF268032_1	
5647	14916439	(AF268032) rhophilin-like protein [Homo sapiens]	2E-14

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
	Į.	gi 7227910 sp O77408 OAR1_LYMST OCTOPAMINE	
		RECEPTOR 1 (OA1) gb AAC61296.1  (U62771)	
5648	7227910	octopamine receptor type 1 [Lymnaea stagnalis]	5.5
		gi 13161815 emb CAC32956.1  (AJ293063) cytochrome	
5649	13161815	oxidase I [Alcochara punctatella]	7.9
	l	gi 13631750 ref XP_010294.3  arylsulfatase E precursor	
5651	13631750	[Homo sapiens]	2E-12
		gi 9181886 gb AAF85678.1 AF266288_4 (AF266288)	
5653	9181886		3.3
		gi 14755042 ref XP_045122.1  KIAA1451 protein	
5654	14755042	[Homo sapiens]	0.000000005
		gi 9408429 gb AAF87294.1  (AF228467) unknown	
5655	9408429	[Letharia vulpina]	2.4
	<b></b>	gi 14530763 cmb CAC42469.1  (AL137139)	
5656	14530763	bA134O15.1 (similar to citrate lyase) [Homo sapiens]	2E-11
	ĺ		
		gi 2496882 sp Q11187 YPD8_CAEEL	
	i	HYPOTHETICAL 99.0 KD PROTEIN C05D11.8 IN	
		CHROMOSOME III gb AAB53828.1  (U00048)	
5661	2496882	C05D11.8 gene product [Caenorhabditis elegans]	4.9
	l <b></b>	gi 1276867 gb AAA97866.1  (U40233) alkane	
5663	1276867	hydroxylase [Stenotrophomonas maltophilia]	8.2
	l	gi 4867907 dbj BAA77721.1  (AB008177) hepatic	
5664	4867907	nuclear factor 1-beta short form [Mus musculus]	3.8
		gi 7292501 gb AAF47904.1  (AE003481) CG11345	
5667	7292501	gene product [Drosophila melanogaster]	3.9
	ł	gi 1346543 sp P49285 ML1A_CHICK MELATONIN	
		RECEPTOR TYPE 1A (MEL-1A-R) (CKA)	
5.000		gb AAA92498.1  (U31820) Mel-1a melatonin receptor	_
5673	1346543	[Gallus gallus]	7
5.075	147017	gi 14731714 ref XP_028009.1  KIAA1563 protein	477.0
5675	14/31/14	[Homo sapiens]	4E-26
		gi 9628467 ref NP_043350.1  putative [Human	
5676	0630463	papillomavirus type 21] gb AAA79398.1  (U31779)	
36/6	9628467	putative [Human papillomavirus type 21] gi 13241978 gb AAK16497.1 AF329199 1 (AF329199)	1.2
5680	12241070		1.0
3080	132419/8	CocoaCrisp [Gallus gallus]	1.9
	l	gi 11466217 ref NP_066540.1  SecY-independent	
		transporter protein [Naegleria gruberi] gb AAG17818.1 AF288092 43 (AF288092) SecY-	
5681	11466217	gb AAG17818.1 AF288092_43 (AF288092) SecY- independent transporter protein [Naegleria gruberi]	4.2
3081	11406217	mucpendent transporter protein [Naegieria gruberi]	4.2

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESS	John Tolling Tolling	1	
NO	N	DESCRIPTION	P VALUE	
		gi 10581459 gb AAG20194.1  (AE005096) enoyl-CoA		
5682	10581459	hydratase; Fad1 [Halobacterium sp. NRC-1]	5	
		gi 4903124 dbj BAA78009.1  (AB027308) nucleotide-		
5684	4903124	binding protein [Plasmid R64]	3.1	
		gi 13242251 ref[NP_077336.1  hairy and enhancer of		
		split 1, (Drosophila) [Rattus norvegicus]		
		sp Q04666 HES1_RAT TRANSCRIPTION FACTOR		
		HES-1 (HAIRY AND ENHANCER OF SPLIT 1)		
İ		(HAIRY-LIKE) (RHL) pir  S36748 transcription factor		
	ļ	HES-1 - rat dbj BAA02682.1  (D13417) HES-1 factor		
	l	[Rattus norvegicus] prf  1905315A HES-1 protein		
5685	13242251	[Rattus norvegicus]	9.4	
		gi 9957880 gb AAG03352.1  (AY005440) 51-kDa		
5689	9957880	antigen [Ehrlichia risticii]	6.9	
		gi 7509017 pir  T33819 hypothetical protein W05F2.7 -		
	2500012	Caenorhabditis elegans gb AAC78217.1  (AF106582)		
5691		Hypothetical protein W05F2.7 [Caenorhabditis elegans]	0.82	
5692	1082343	gi 1082343 pir  S50832 atrophin-1 - human	0.05	
		1115055001 1101 010010 11/17 151010		
5696	11505620	gi 11595629 cmb CAC18249.1  (AL451018) conserved hypothetical protein [Neurospora crassa]	2.6	
3090			3.6	
		gi 11465898 ref NP_066447.1  ATP synthase F0 subunit 6 [Ochromonas danica] gb AAG18413.1 AF287134 38		
		(AF287134) ATP synthase F0 subunit 6 [Ochromonas		
5698	11465898		6,6	
3098	11403898	danicaj	0.0	
	ļ			
		gi 13384736 ref NP_084514.1  dynein, cytoplasmic, heavy chain 1; dynein heavy chain, retrograde transport		
		Incavy chain 1; dyncin heavy chain, retrograde transport   Mus musculus   sp   Q9JHU4   DYHC MOUSE DYNEIN		
		HEAVY CHAIN, CYTOSOLIC (DYHC)		
		(CYTOPLASMIC DYNEIN HEAVY CHAIN)		
		gb AAF91078.1  (AY004877) cytoplasmic dynein heavy		
5699	13384736	chain [Mus musculus]	3E-12	
		gi 11431299 ref XP 007961.1  phosphorylase kinase,		
5701	11431299	beta [Homo sapiens]	8.8	
		gi 7486830 pir  T04917 hypothetical protein T10I14.190		
		- Arabidopsis thaliana emb CAA16786.1  (AL021712)	-	
		putative protein [Arabidopsis thaliana]		
		emb CAB79191.1  (AL161557) putative protein		
5704	7486830	[Arabidopsis thaliana]	0.4	

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS	The state of the s	1 Totolias)
NO	N	DESCRIPTION	P VALUE
		gi 7470729 pir  S75327 lysostaphin - Synechocystis sp.	
	ŀ	(strain PCC 6803) dbj BAA17241.1  (D90904)	
5707	7470729	lysostaphin [Synechocystis sp. PCC 6803]	8.8
		gi 7330072 gb AAF60061.1 AF210726 82 (AF210726)	
5708	7330072	ORFRU4-R [Macaca mulatta rhadinovirus 26-95]	0.24
		gi 14774911 ref XP 027365.1  ADP-ribosylation factor-	
5711	14774911	like 6 interacting protein [Homo sapiens]	1E-11
		gi 14721018 ref XP 051562.1  similar to agrin (H.	
5712	14721018	sapiens) [Homo sapiens]	0.14
		gi 6324014 ref NP 014084.1  F1FO ATPasc assembly	
1	ł	protein; Atpl1p [Saccharomyces cerevisiae]	
	1	sp[P32453]ATPS YEAST ATP11 PROTEIN	
	ì	PRECURSOR pir  S51297 ATP11 protein - yeast	
Ì		(Saccharomyces cerevisiae) gb AAA34447.1  (M87006)	
		ATP11 protein [Saccharomyces cerevisiae]	
		cmb CAA86381.1  (Z46259) ATP11 [Saccharomyces	
l	ŀ	cerevisiae] emb CAA96245.1  (Z71591) ORF YNL315c	
5716	6324014	[Saccharomyces cerevisiae]	4.1
		gi 12861848 dbj BAB32292.1  (AK021073) putative	
5718	12861848	[Mus musculus]	0.07
5719	14744130	gi 14744130 ref XP_045895.1  57406 [Homo sapiens]	1.4
		gi 11612206 gb AAG37299.1  (AY009937) unknown	
5722	11612206	[Sinorhizobium fredii]	7.4
		gi 4494967 gb AAD21389.1  (AF083501) R11 [Macaca	
		mulatta rhadinovirus 17577]	
		gb AAF60041.1 AF210726_62 (AF210726) vIRF	
5723	4494967	[Macaca mulatta rhadinovirus 26-95]	3.3
	ŀ		
	1	gi 12644379 sp Q02099 RAD3_SCHPO DNA REPAIR	
		PROTEIN RAD3 pir  T39911 rad3 checkpoint protein -	
		fission yeast (Schizosaccharomyces pombe)	
		emb CAB40165.1  (AL049558) rad3 checkpoint protein	
5727	12644379	[Schizosaccharomyces pombe]	9.3
		gi 7492633 pir  T40241 probable guanine nucleotide	
		exchange factor - fission yeast (Schizosaccharomyces	
		pombe) emb CAB58155.1  (AL121815) putative	
		guanine nucleotide exchange factor	
5732	7492633	[Schizosaccharomyces pombe]	8.2
		gi 10047255 dbj BAB13416.1  (AB046810) KIAA1590	
5737	10047255	protein [Homo sapiens]	0.00000002

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 2498204 sp Q05682 CALD_HUMAN	
		CALDESMON (CDM) pir  JH0628 caldesmon - human	
5742	2498204	gb AAA58419.1  (M83216) caldesmon [Homo sapiens]	8E-23
		gi 11037059 ref]NP_036540.1  protein kinase C binding	
		protein 1 [Homo sapiens] gb AAF71262.1  (AF233453)	
5744	11037059	RACK-like protein PRKCBP1 [Homo sapiens]	0.00000001
		gi 14575679 gb AAK68690.1 AF156100_1 (AF156100)	
5746	14575679	hemicentin [Homo sapiens]	1.9
		gi 11466210 ref NP_066533.1  NADH dehydrogenase	
		subunit 2 [Naegleria gruberi]	
		gb AAG17811.1 AF288092_36 (AF288092) NADH	
5748	11466210	dehydrogenase subunit 2 [Naegleria gruberi]	8.2
		gi 8096340 dbj BAA95898.1  (AB036737) RERE	
5750	8096340	[Homo sapiens]	0.5
		gi 14275752 emb CAC40032.1  (AJ310844) P-type	
5751	14275752	ATPase [Hordeum vulgare]	0.53
		gi 5834582 emb CAB55313.1  (AJ132948) rfg7 protein	
5752	5834582	[Homo sapiens]	1.7
		gi 13877921 gb AAK44038.1 AF370223_1 (AF370223)	
5754	13877921	unknown protein [Arabidopsis thaliana]	4
		gi 5869818 emb CAB55575.1  (AJ249395) NADH-	
		ubiquinone oxidoreductase subunit 6 [Globodera	
5755	5869818	pallida]	4.2
	7000000	gi 7023033 dbj BAA91809.1  (AK001649) unnamed	100
5756	7023033	protein product [Homo sapiens]	e-102
	1.4500.665	gi 14729667 ref XP_029101.1  KIAA0947 protein	
5757	14729667	[Homo sapiens]	0.22
		gi 14726213 ref XP_010556.3  PR domain containing 16	
5758	14726213	[Homo sapiens]	0.87

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 4757900 ref NP_004334.1  calreticulin precursor; Sicca syndrome antigen A (autoantigen Ro; calreticulin); autoantigen Ro [Homo sapiens] ref XP_03202.1  calreticulin precursor [Homo sapiens] ref XP_03202.1  calreticulin precursor [Homo sapiens] ref XP_093055.3  calreticulin precursor [Homo sapiens] ref XP_09055.3  calreticulin precursor [Homo sapiens] ref XP_09055.3  calreticulin precursor [Homo sapiens] ref XP_093202.1  calreticulin precursor [Homo sapiens] sp P2797 CRTC_HUMAN CALRETICULIN PRECURSOR (CRP55) (CALREGULIN) (HACBP) (ERP60) (32 KDA RIBONUCLEOPROTEIN AUTOANTIGEN RO/SS-A) pir A37047 calreticulin precursor - human gb AAA51916.1  (M\$4739) calreticulin [Homo sapiens] gb AAA5682.1  (M32294) Ro ribonucleoprotein autoantigen (Ro/SS-A) precursor [Homo sapiens] gb AAB51176.1  (AD000092) calreticulin [Homo sapiens] gb AAB510500 (BC002500) calreticulin	. 11000
5761	4757900	[Homo sapiens] gb AAH07911.1 AAH07911	5E-16
		gi 14714688 gb AAH10485.1 AAH10485 (BC010485)	
5762	14714688	Unknown (protein for MGC:7224) [Mus musculus]	1.1
		gi 14751380 ref XP 041291.1  cAMP response element-	
5765	14751380	binding protein CRE-BPa [Homo sapiens]	5E-14
		gi 2105193 gb AAB57925.1  (U86889) hypothetical	
5767	2105193	protein [Molluscum contagiosum virus subtype 1]	7.8
5771		gi 13278418 gb AAH04019.1 AAH04019 (BC004019) Similar to mesoderm specific transcript [Mus musculus]	0.94
		gi 14720481 rcf XP_048811.1  hypothetical protein	
		FLJ22116 [Homo sapiens] rcf[XP_048810.1]	
5772	14720481	hypothetical protein FLJ22116 [Homo sapiens]	2E-23

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
	ł		
		gi 6321548 ref NP_011625.1  Ygr110wp	
		[Saccharomyces cerevisiae] sp P53264 YG2V_YEAST	
J		HYPOTHETICAL 52.0 KD PROTEIN IN CLB6-	
ì	ì	SHY1 INTERGENIC REGION pir S64418	
ļ		hypothetical protein YGR110w - yeast (Saccharomyces	
5773	(221540	cerevisiae) emb CAA97118.1  (Z72895) ORF	
3113	0321348	YGR110w [Saccharomyces cerevisiae]	2.4
5776	10050040	gi 12853842 dbj BAB29864.1  (AK015480) putative  Mus musculus	
5//6	12853842	[Mus musculus]	5.6
		gi 13384736 rcf NP_084514.1  dyncin, cytoplasmic,	
		heavy chain 1; dynein heavy chain, retrograde transport	
		[Mus musculus] sp Q9JHU4 DYHC_MOUSE DYNEIN	
l		HEAVY CHAIN, CYTOSOLIC (DYHC)	
		(CYTOPLASMIC DYNEIN HEAVY CHAIN)	
5778	12204526	gb AAF91078.1  (AY004877) cytoplasmic dynein heavy chain [Mus musculus]	25.02
3//6	13364730	gi 7498705 pir T20640 hypothetical protein F09C3.3 -	2E-92
5781	7498705	Caenorhabditis elegans	9.7
3701	1420103	gi 11356402 pir  T44074 hypothetical protein [imported]	9.7
5782	11356402	- Staphylococcus aureus (fragment)	5,5
5702	11550102	gi 10726396 gb AAF54288,2  (AE003680) CG11773	3.3
5783	10726396		4.1
	11120010	gi 11935116 gb AAG41977.1 AF311942 1 (AF311942)	
5784	11935116	ethylene receptor [Carica papaya]	7.5
		gi 4456467 cmb CAB37294.1  (AJ011001) TM7XN1	- 1.0
5787	4456467	protein [Homo sapiens]	3.8
		gi 13676779 gb AAK38272.1 AF330197 1 (AF330197)	
5788	13676779	Arkadia [Mus musculus]	2.2
		gi 13676779 gb AAK38272.1 AF330197_1 (AF330197)	
5789	13676779	Arkadia [Mus musculus]	2.3
		gi 11024704 ref NP_061956.1  hypothetical protein	
		FLJ11219 [Homo sapiens] dbj BAA92074.1	
5795	11024704	(AK002081) unnamed protein product [Homo sapiens]	8E-23
1			
1		gi 13606086 gb AAK32948.1  (U41014) Hypothetical	
5797	13606086	protein C06G1.1 [Cacnorhabditis elegans]	8.3
		gi 10728595 gb AAF52302.2  (AE003611) CG9011	_
5798	10728595	gene product [Drosophila melanogaster]	8E-21
5700	100 45055	gi 12847975 dbj BAB27780.1  (AK011690) putative	27.44
5799	12847975	[Mus musculus]	3E-44

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 7243247 dbj BAA92671.1  (AB037854) KIAA1433	
5801	7243247	protein [Homo sapiens]	3E-84
		gi 10728595 gb AAF52302.2  (AE003611) CG9011	
5803	10728595	gene product [Drosophila melanogaster]	8E-21
		gi 1354082 gb AAB37237.1  (U47033) polyprotein	
5804	1354082	[bean yellow mosaic virus]	6,8
		gi 11354734 pir  C82043 conserved hypothetical protein	
		VC2703 [imported] - Vibrio cholerae (group O1 strain	
		N16961) gb AAF95843.1  (AE004336) conserved	
5808	11354734	hypothetical protein [Vibrio cholerae]	4.9
		gi 13272353 gb AAK17116.1 AF291051_13	
		(AF291051) transketolase [Candidatus Carsonella	
5809	13272353		8.9
		gi 10728595 gb AAF52302.2  (AE003611) CG9011	
5810	10728595	gene product [Drosophila melanogaster]	9E-21
	1	gi 7340927 dbj BAA92999.1  (AP001550) Similar to	
1	ł	Schizosaccharomyces pombe chromosome I cosmid	
		c1D4; hypothetical protein &SPAC1D4_10 (Q10155)	
5818	7340927	[Oryza sativa]	5.6
		gi 4115911 gb AAD03423.1  (U78517) cAMP-regulated	
		guanine nucleotide exchange factor II [Rattus	
5821	4115911	norvegicus]	3E-11
1		gi 7498336 pir  T15086 hypothetical protein E03D2.4 -	
1		Caenorhabditis clegans gb AAB94164.1  (AF039036)	
5822	7498336	Hypothetical protein E03D2.4 [Caenorhabditis elegans]	7.1
		gt 7492270 pir  T40528 palmitoyl-protein thioesterase	
		precursor - fission yeast (Schizosaccharomyces pombe)	
1	ł	emb CAA19178.1  (AL023634) palmitoyl-protein	
5825	7492270	thioesterase precursor [Schizosaccharomyces pombe]	1.2
1		gi 7497138 pir  T30158 hypothetical protein C37A2.6 -	
	1	Cacnorhabditis clegans gb AAB52451.1  (U97194)	·
5827	7497138	Hypothetical protein C37A2.6 [Caenorhabditis elegans]	5.4
l	ĺ	gi 7469981 pir  S74598 hypothetical protein sll1040 -	
		Synechocystis sp. (strain PCC 6803) dbj BAA16750.1	
		(D90900) ORF_ID:sl11040~unknown protein	
5830	7469981	[Synechocystis sp. PCC 6803]	9.4

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	Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 10177727 dbj BAB10973.1  (AB015471)	
		gene_id:K9B18.6~unknown protein [Arabidopsis	
5831	10177727		2.6
		gi 2231323 gb AAB61999.1  (U77045) bactinecin 6	
5834	2231323	[Ovis aries]	6.9
		' i i i i i i i i i i i i i i i i i i i	
		gi 7495511 pir  T18990 hypothetical protein C06B8.4 -	
Į.		Caenorhabditis elegans emb CAB03847.1  (Z81463)	
		predicted using Genefinder~Similarity to C.elegans	
		olfactory receptor ODR-10 (TR:Q17376), contains	
i		similarity to Pfam domain: PF01461 (7TM	1
		chemoreceptor), Score=-92.4, E-value=4.9e-10, N=1	
5838	7495511	[Caenorhabditis elegans]	0.84
		gi 14722156 ref XP_001418.4  centromere protein F	
5839	14722156	(350/400kD) [Homo sapiens]	2
		gi 14590902 ref[NP_142975.1  hypothetical protein	
		[Pyrococcus horikoshii] pir  E71100 hypothetical protein	
		PH1064 - Pyrococcus horikoshii dbj BAA30163.1	
		(AP000004) 718aa long hypothetical protein	
5841	14590902	[Pyrococcus horikoshii]	3.5
		gi 10173304 dbj BAB04409.1  (AP001509) transposase	
5844	10173304	(04) [Bacillus halodurans]	2.2
		gi 7510388 pir  T27298 hypothetical protein Y68A4A.7 -	
1		Caenorhabditis elegans emb CAA16418.1  (AL021503)	
		predicted using Genefinder~contains similarity to Pfam	
		domain: PF01604 (7TM chemoreceptor), Score=-48.3,	
5845	7510388	E-value=7.1e-07, N=1 [Caenorhabditis elegans]	2.9
		gi 13812127 ref NP_113254.1  hypothetical protein	
		[Guillardia theta] gb AAK39814.1 AF165818_22	
5851	13812127	(AF165818) hypothetical protein [Guillardia theta]	0.92
		gi 6016842 dbj BAA85182.1  (AB033168) nuclear	
5852	6016842	protein ZAP [Mus musculus]	3.3
		gi 8922744 ref[NP_060730.1  hypothetical protein	
		FLJ10891 [Homo sapiens] dbj BAA91884.1	
5854	8922744	(AK001753) unnamed protein product [Homo sapiens]	8E-69

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 118965 sp P23098 DYHC_TRIGR DYNEIN BETA	
[		CHAIN, CILIARY pir  S17653 dynein beta heavy	
		chain, ciliary - sea urchin (Tripneustes gratilla)	
		emb CAA42170.1  (X59603) Beta heavy chain of outer-	
		arm axonemal dynein ATPase [Tripneustes gratilla]	
	l	prf∥1714372A dyncin:SUBUNIT=beta heavy chain	
5856	118965	[Tripncustes gratilla]	0.0000001
		gi 10640323 emb CAC12137.1  (AL445066) conserved	
	İ	hypothetical membrane protein [Thermoplasma	
5858	10640323	acidophilum]	1.3
		gi 9964574 rcf NP_065042.1  AMV260 [Amsacta	
		moorei entomopoxvirus]	
		gb AAG02966.1 AF250284_260 (AF250284) AMV260	
5859	9964574		4
		gi 5360226 dbj BAA36472.1  (AB015177) F0-ATPase	
5861	5360226	subunit 6 [Beta vulgaris]	3
i l		gi 7327641 gb AAF45040.2  (AF146609) putative	
5868	7327641	modification methyltransferase [Aeromonas hydrophila]	2.8
		gi 5031969 ref NP_005758.1  purinergic receptor	
		(family A group 5) [Homo sapiens]	
		sp P43657 P2Y5_HUMAN P2Y PURINOCEPTOR 5	
		(P2Y5) (PURINERGIC RECEPTOR 5) (RB INTRON	
		ENCODED G-PROTEIN COUPLED RECEPTOR)	
		pir  T09508 intron 17 purinergic receptor P2Y5 - human	
		gb AAB62190.1  (AF000546) purinergic receptor P2Y5	
5871	5031969	[Homo sapiens]	3.2
		gi 92972 pir  S04757 NADH dehydrogenase	
5872	92972	(ubiquinone) (EC 1.6.5.3) chain 5 - rat mitochondrion	9.8
		gi 13540669 ref NP_110480.1  linker for activation of T	
		cells [Rattus norvegicus] sp O70601 LAT_RAT	
		LINKER FOR ACTIVATION OF T CELLS (36 KDA	
		PHOSPHO-TYROSINE ADAPTOR PROTEIN)	
	l	(PP36) (P36-38) emb CAA04577.1  (AJ001184) 36 kDa	
5876	13540669	phospho-tyrosine [Rattus norvegicus]	0.4

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 7510482 pir T27406 hypothetical protein Y75B8A.24	
		- Caenorhabditis elegans cmb CAA22108.1	
		(AL033514) contains similarity to Pfam domain:	
	l	PF00454 (Phosphatidylinositol 3- and 4-kinases),	
		Score=392.3, E-value=1.6e-114, N=2; PF00613	
		(Phosphoinositide 3-kinase family, accessory domain	
		(PIK domain)), Score=76.1, E-value=2.4e-19,	
5877	7510482	N=1~cDNA EST yk2>	0.32
		gi 11284162 pir  H81077 hypothetical protein NMB1490	
		[imported] - Neisseria meningitidis (group B strain	
		MD58) gb AAF41846.1  (AE002498) hypothetical	
5881	11284162	protein [Neisseria meningitidis MC58]	4.2
		gi 4503801 ref NP_003893.1  far upstream element-	
	İ	binding protein; far upstream element binding protein;	
		FUSE-binding protein [Homo sapiens] pir  A53184 myc	
	l	far upstream element-binding protein - human	
5882	4503801	gb AAA17976.1  (U05040) FUSE binding protein	0.4
5882	4503801		0.4
5883	9929953	gi 9929953 dbj BAB12133.1  (AB047609) hypothetical protein [Macaca fascicularis]	9.7
3003	9929933	gi 13631383 ref XP 010272.2  retinoblastoma-binding	9.1
		protein 7 [Homo sapiens] ref[XP_045112.1]	
5884	13631383	retinoblastoma-binding protein 7 [Homo sapiens]	7.3
3004	13031363	gi 7662084 ref NP 055474.1  KIAA0377 gene product	7.3
		[Homo sapiens] dbj BAA20831.1  (AB002375)	
5886	7662084	KIAA0377 [Homo sapiens]	2E-44
3000	7002004	Kirrios // [riono sapions]	25-11
		gi 4507023 ref[NP 003031.1  solute carrier family 4,	
		anion exchanger, member 2 (erythrocyte membrane	
		protein band 3-like 1) [Homo sapiens]	
		sp[P04920]B3A2 HUMAN ANION EXCHANGE	
		PROTEIN 2 (NON-ERYTHROID BAND 3-LIKE	
	l	PROTEIN) (BND3L) pir S21086 anion exchange	
		protein 2 - human emb CAA44067.1  (X62137) anion	
5888	4507023	exchange protein 2 (AE2) [Homo sapiens]	9.1

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID NO	ACCESS N	DESCRIPTION	P VALUE
		gi 6321764 ref NP_011840.1  Yhl023cp	
		[Saccharomyces cerevisiae] sp P38742 YHC3_YEAST	
		HYPOTHETICAL 130.0 KD PROTEIN IN SNF6-	
	İ	SPO11 INTERGENIC REGION pir \$46837	
		hypothetical protein YHL023c - yeast (Saccharomyces cerevisiae) gb AAB65076.1  (U11582) No definition line	
5892	6321764	found [Saccharomyces cerevisiae]	7.4
3692	0321704	ionid [Saccharoniyees cerevisiae]	7.4
		gi 7507827 pir  T16876 hypothetical protein T14E8.3 -	
		Caenorhabditis elegans gb AAA82386.1  (U41036)	
5894	7507827	Hypothetical protein T14E8.3 [Cacnorhabditis elegans]	2.4
		gi 119714 sp P13983 EXTN TOBAC EXTENSIN	
		PRECURSOR (CELL WALL HYDROXYPROLINE-	
		RICH GLYCOPROTEIN) pir  S06733 hydroxyproline-	
		rich glycoprotein precursor - common tobacco	
		emb CAA32090.1  (X13885) extensin (AA 1-620)	
5896	119714	[Nicotiana tabacum]	2.4
		gi 13540669 rcf NP_110480.1  linker for activation of T	
		cells [Rattus norvegicus] sp O70601 LAT_RAT	
		LINKER FOR ACTIVATION OF T CELLS (36 KDA	
		PHOSPHO-TYROSINE ADAPTOR PROTEIN)	
5897	125 406 60	(PP36) (P36-38) emb CAA04577.1  (AJ001184) 36 kDa phospho-tyrosine [Rattus norvegicus]	0.4
3097	13340009	gi 14787176 gb AAG54083,1  (AY017475) CSMD1	0.4
5898	14787176	[Mus musculus]	0.5
		gi 14192869 gb AAK55774,1 AC079038 8	
5899	14192869	(AC079038) Putative polyprotein [Oryza sativa]	1.9
		gi 11260604 pir  G82485 acetate kinase VCA0235	
		[imported] - Vibrio cholerae (group O1 strain N16961)	
5000	1100000	gb AAF96146.1  (AE004363) acetate kinase [Vibrio	
5900	11260604	choleraej	5.4
		gi 5688864 dbj BAA82706.1  (AB030586) amino acid	
5914	5688864	transporter-like protein 1 [Arabidopsis thaliana]	0.74
		ail4571551 abla AV 64511    (AV026002) 3	
5915	14571551	gi 14571551 gb AAK64511.1  (AY036902) degenerative spermatocyte-like protein RDES [Rattus norvegicus]	7.1
	2.13.11331	gi 11496539 ref NP 044549.1  ribosomal protein S3	***
		Toxoplasma condiil cblAAD41136.1lU87145 5	
5917	11496539	(U87145) ribosomal protein S3 [Toxoplasma gondii]	2.4

Table 3B Nearest Neighbor (BlastX vs. Non-Redundant Proteins)								
SEQ ID ACCESS								
NO	N	DESCRIPTION	P VALUE					
		gi 12275264 cmb CAC22282.1  (AJ303456) WASP						
5919	12275264	interacting protein [Rattus norvegicus]	0.29					
		gi 7158837 gb AAF37557.1 AF214067_1 (AF214067)						
5920	7158837	serine-repeat antigen protein [Plasmodium falciparum]	3.8					
		gi 7327641 gb AAF45040.2  (AF146609) putative						
5922	7327641	modification methyltransferase [Aeromonas hydrophila]	3.2					
		gi 4321845 gb AAD15841.1  (AF064782) unknown						
5934	4321845	[Mus musculus]	4E-59					
		gi 6562754 emb CAB62893.1  (AL035475) hypothetical						
5938	6562754		0.69					
		gi 13814462 gb AAK41504.1  (AE006741) ABC						
		transporter, ATP binding protein [Sulfolobus						
5939	13814462	solfataricus]	2.3					
		gi 5802786 gb AAD51779.1  (AF124511) BVES						
5941	5802786	[Gallus gallus]	1.7					
		gi 5596342 dbj BAA82602.1  (AB026825) sALK-2						
5943	5596342		7.2					
	1	gi 9802527 gb AAF99729.1 AC004557_8 (AC004557)						
5946	9802527		6.8					
		gi 3820854 emb CAA10852.1  (AJ222582) maturase-						
5949	3820854	like protein [Euglena granulata]	4.2					
	ľ	gi 7476228 pir  A70905 hypothetical protein Rv0174 -						
		Mycobacterium tuberculosis (strain H37RV)						
		emb CAB09741.1  (Z97050) hypothetical protein						
5951	7476228	Rv0174 [Mycobacterium tuberculosis]	0.46					
	l							
		gi 7159336 gb AAF37725.1 AF238235_1 (AF238235)						
5953	7159336	diaphanous protein [Entamoeba histolytica]	0.24					
		gi 7294100 gb AAF49454.1  (AE003527) CG12243						
5954	7294100	gene product [Drosophila melanogaster]	4.6					
		gi 2689578 gb AAB91357.1  (U96421) cytochrome b						
5956	2689578		9.4					
		gi 11071788 emb CAC14632.1  (AL449144)						
5957	11071788	hypothetical protein P214.26 [Leishmania major]	5.7					

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		110101112)
NO	N	DESCRIPTION	P VALUE
		gi 9631305 ref NP 048116.1  ORF MSV045	
		hypothetical protein [Melanoplus sanguinipes	
		entomopoxvirus] pir T28206 hypothetical protein	
		ORF45 - Melanoplus sanguinipes entomopoxvirus	
		gb AAC97618.1  (AF063866) ORF MSV045	
		hypothetical protein [Melanoplus sanguinipes	
5958	9631305	entomopoxvirus]	3
		gi 14750752 ref XP_031441.1  phosphodiesterase 8A	
5959	14750752	[Homo sapiens]	0.47
		gi 4996347 dbj BAA78416.1  (AB021177) complement	
5962	4996347	B/C2-A2 [Cyprinus carpio]	1.7
		gi 7510076 pir  T31613 hypothetical protein Y50E8A.i -	
5964	7510076		7.1
		gi 6958206 gb AAF32493,1 AF093132_1 (AF093132)	
		kexin-like protease KEX1 [Pneumocystis carinii f. sp.	
5965	6958206	muris]	4.5
		gi 13542701 gb AAH05557.1 AAH05557 (BC005557)	
5966	13542701	Unknown (protein for MGC:7062) [Mus musculus]	0.00003
		gi 12852967 dbj BAB29595.1  (AK014872) putative	
5967	12852967	[Mus musculus]	3.2
		gi 6449214 gb AAF08856.1 AF194824_1 (AF194824)	
5969	6449214	NADH dehydrogenase [Aptenia cordifolia]	0.84
		gi 7492532 pir  T39653 probable DNA repair and	
		recombination protein - fission yeast	
		(Schizosaccharomyces pombe) emb CAA21300.1	
5971	7492532	(AL031856) putative DNA repair and recombination	
39/1	1492532	protein [Schizosaccharomyces pombe]	8.4
		gi 1171089 sp P10243 MYBA_HUMAN MYB- RELATED PROTEIN A (A-MYB) pir  S03423	
5972	1171089	transforming protein A-myb - human	2E-28
3912	11/1089	gi 4589488 dbi BAA76772.1  (AB023145) KIAA0928	ZE-28
5973	4589488	gi 4389488 doj BAA/6//2.1  (AB023143) KIAA0928 protein [Homo sapiens]	9.8
3913	4509400	gi 6686326 sp P77589 MHPT ECOLI PUTATIVE 3-	9.0
		HYDROXYPHENYLPROPIONIC ACID	
5974	6686326	TRANSPORTER	1.8
37/4	0000320	gi 7298382 gb AAF53607.1  (AE003655) CG15141	1.0
5976	7298382	gene product [Drosophila melanogaster]	5.7
3710	1230302	Some Product [D1020hima incianogaster]	J.1

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		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		,
NO.	N	DESCRIPTION	P VALUE
		gi 7477896 pir  D70595 probable ATP-dependent RNA	
		helicase - Mycobacterium tuberculosis (strain H37RV)	
		emb CAB08305.1  (Z95120) rhlE [Mycobacterium	
		tuberculosis] gb AAK47649.1  (AE007143) ATP-	
		dependent RNA helicase DeaD [Mycobacterium	
5979	7477896	tuberculosis CDC1551]	4.7
		gi 10726674 gb AAF55880.2  (AE003734) CG16791	
5980	10726674	gene product [Drosophila melanogaster]	5,5
		gi 10303299 emb CAC10094.1  (AL442164) related to	
5983	10303299	SEN1 protein [Neurospora crassa]	9.4
		gi 14625441 dbj BAB61903.1  (AB053446) KIAA1773	
5984	14625441	protein [Homo sapiens]	2.4
		gi 2496701 sp P55552 Y4LL_RHISN	
		HYPOTHETICAL 91.8 KD PROTEIN Y4LL	
****		gb AAB91764.1  (AE000083) Y4lL [Rhizobium sp.	
5985	2496701	NGR234]	44
		gi 13652830 ref XP_017068.1  65851 [Homo sapiens]	
		ref XP_039450.1  similar to NONHISTONE	
5988	10050000	CHROMOSOMAL PROTEIN HMG-14 (H. sapiens)	0.00001
3988	13652830	[Homo sapiens]	0,00001
		gi 2098719 gb AAB57675.1  (U85709) putative fimbrial-	
5989	2098719	associated protein [Actinomyces naeslundii]	5.4
3769	2090/19	gil1142588gblAAA84740.1l (U05313) CR3	3.4
5990	1142588	[Trypanosoma brucei]	0.02
3770	1142500	gi 11283273 pir  A81658 hypothetical protein TC0845	0.02
		[imported] - Chlamydia muridarum (strain Nigg)	
		gb AAF39643.1  (AE002351) hypothetical protein	
5991	11283273	[Chlamydia muridarum]	7.2
		gi 13507856 ref[NP 109805.1  ribosomal protein L20	
		[Mycoplasma pneumoniae] sp[P78023]RL20 MYCPN	
		50S RIBOSOMAL PROTEIN L20 pir S73363	
		ribosomal protein L20 - Mycoplasma pneumoniae	
		(strain ATCC 29342) gb AAG34734.1 AE000004 3	
		(AE000004) ribosomal protein L20 [Mycoplasma	
5992	13507856	pneumoniae]	0.84
		gi 7506563 pir  T24113 hypothetical protein R10D12.3	
		Caenorhabditis elegans emb CAB03243.1  (Z81109)	
		predicted using Genefinder-similar to G-protein coupled	
5997	7506563	receptor [Caenorhabditis elegans]	6.3
		gi 7510771 pir  T29919 hypothetical protein ZC449.5 -	
5998	7510771	Caenorhabditis elegans	3.6

		Table 3B Nearest Neighbor (BlastX vs. Non-Redundant	Proteins)
SEQ ID	ACCESS		
NO	N	DESCRIPTION	P VALUE
		gi 14755125 ref XP_006601.4  hypothetical protein	
6001	14755125	FLJ10659 [Homo sapiens]	3E-96
6005	7508360	gi[7508360]pir[[T25220 hypothetical protein T24B8.4 - Caenorhabditis elegans emb[CAA92756.1] (Z68338) predicted using Genefinder-contains similarity to Pfam domain: PF02205 (Wiskott Aldrich syndrome homology region 2), Score=43.6, E-value=1.5e-09, N=2~cDNA EST yk96d7.5 comes from this geno~cDNA EST yk96d7.3 comes from this geno~cDNA EST yk76d1>	1
		gi 4960210 gb AAD34644.1 AF154112_1 (AF154112)	
6006	4960210	transcription co-repressor Sin3 [Xenopus laevis]	7

Table 4					
SEQ ID NO	SEQ NAME	CLUSTER	PROFILE NAME	DIR	SCORE
6	2102.B18.gz43_275316	558147	Ets_Cterm	for	19.58
9	2103.M06.gz43_275519	377696	protkinase	for	20.71
38	2153,K14,gz43_278937	372952	Dead_box_helic	for	172.21
39	2154.M04.gz43_279163	377696	protkinase	for	20.71
61	2165.H06.gz43_280342	393635	zf-c2h2	for	33.96
69	2166,J11.gz43_281368	377696	protkinase	for	20,71
108	2118.A09.gz43_307025	446397	bzip	for	19.15
117	2131.113.gz43_308085	34071	wd40	for	37.45
118	2131.B14.gz43_308094	221686	protkinase	for	33,14
228	1573.F18.gz43_208848	639849	PH	for	42,77
229	1573.K19.gz43_208869	486238	protkinase	rev	45.41
415	1585.G22.gz43 210545	412416	Dead box helic	for	49,67
445	1587,B06,gz43 211440	446984	ANK	rev	23.12
486	1597.G06.gz43_212233	639593	defensins	rev	18.27
487	1597.J06.gz43_212236	557975	ANK	for	35.63
502	1597.F18.gz43_212424	596882	zf-c2h2	rev	18.13
700	1694.M19.gz43 214375	425923	zf-c2h2	for	32.76
847	1706,P07.gz43_216138	639901	zf-c2h2	for	19.43
877	1707.J02.gz43 216453	550237	zf-cech	for	26.74
1511	1755,P24.gz43 223395	606129	rvt	for	37.6
1714	1790.C14.gz43 226997	727150	bzip	for	24.2
2034	1828.J19.gz43_232472	728303	zf-c2h2	rev	18.19
2038	1828.P21.gz43_232510	509678	Retvir_asp_protease	for	28.5
2054	1838.N05,gz43 233020	481614	zf-c2h2	for	18.52
2514	1888.O06.gz43_233020	451764	rvt	for	49.99
2973		499700	7tm 1		73.7
	1924.H18.gz43_245579		ANK	rev	
3013	1935,E18,gz43_246500	490805		rev	28.74
3140	1981.O19.gz43_248062	558949	zf-c3hc4	rev	19.16
3403	1958.N12.gz43_250647	556308	zf-c2h2	for	40,77
3524	1923.M22.gz43_252963	562603	zf-c2h2	rey	42.42
3653	1995.C03.gz43_256117	562152	zf-c2h2	rev	18.97
3689	1995,P13.gz43_256290	562989	EGF	rev	19.4
3723	1995.B24.gz43_256452	556632	zf-c2h2	rev	20.64
3814	2007.F09.gz43_257778	560652	zf-c2hc	rev	21.49
3931	2008.F18.gz43_258308	550497	bzip	for	20.27
4151	1669,G11,gz43_260853	503275	protkinase	rev	43.25
4356	1682.O17.gz43_262495	450211	bzip	rev	26.06
4373	1682.F21.gz43_262550	546740	EFhand	rev	18.72
4688	2018.K14.gz43_264760	432970	zf-c2h2	for	48.43
4979	2041.C09.gz43_266976	556632	zf-c2h2	rev	20.88
5467	2067.I20.gz43_271090	551617	7tm_1	rev	19.77
5508	2068.F14.gz43_271375	561707	7tm_1	rev	24.27
5522	2068.D17.gz43_271421	554774	tgf-beta	for	18.24
5756	2176.J17.gz43_281945	412416	Dead_box_helic	for	37.64
6001	1561,C22,gz43_314731	447072	PH	for	31.95

Table 5						
		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
6	558147	12,13	3	12	-1	3.87
11	402353	12.13	13	4	3.36	-1
14	884	03,04	51	20	2,49	-1
14	884	15,17	1	10	-1	9.32
14	884	27,28	15	3	5,41	-1
14	884	28,29	3	21	-l	5.62
22	427571	08,09	6	0	8,38	-1
28	6342	30,31	0	7	-1	6.29
29	387530	30,31	2	24	-1	10.79
52	24210	08,09	2	13	-1	4.65
52	24210	21,22	0	11	-1	11.18
52	24210	23,24	1	9	-1	8.91
52	24210	25,26	12	2	5.79	-1
60	1300	25,26	6	0	5.79	-1
64	376808	08,09	0	8	-1	5,72
65	6342	30,31	0	7	-1	6.29
67	24210	08,09	2	13	-1	4.65
67	24210	21,22	0	11	-1	11.18
. 67	24210	23,24	1	9	-1	8.91
67	24210	25,26	12	2	5.79	-1
68	24210	08,09	2	13	-1	4,65
68	24210	21,22	0	11	-1	11.18
68	24210	23,24	1	9	-1	8.91
68	24210	25,26	12	2	5,79	-1
71	185432	03,04	0	26	-1_	26,65
71	185432	08,09	10	33	-1	2.36
75	418763	15,16	15	3	5,28	-1
75	418763	15,17	15	2	8.05	-1
75	418763	27,29	11	0	14.84	-1
75	418763	28,29	5	0	6,23	-1
79	649035	27,28	0	7	-1	6.46
79	649035	28,29	7	0	8.72	-1
92	186594	25,26	12	0	11,58	-1
93	218904	08,09	6	1	8,38	-1
99	535955	15,17	1	10	-1	9.32
99	535955	16,17	0	10	-1	9.85
112	48238	08,09	6	1	8,38	-1
113	226324	15,17	0	- 8	-1	7.45
113	226324	16,17	0	8	-1	7,88
113	226324	27,29	5	0	6.75	-1
115	62016	16,17	0	6	-1	5.91
116	48238	08,09	6	1	8,38	-1
121	20453	25,26	3	13	-1	4.49
123	37805	08,09	10	0	13.97	-1
125	48238	08,09	6	1	8.38	-1
136	120049	15,17	10	2	5.37	-1

		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
140	446829	27,29	78	44	2.39	-1
147	147196	28,29	16	46	-l	2.31
147	147196	30,31	8	30	-1	3,37
153	446614	16,17	. 7	0	7.11	-1
160	463487	15,17	8	1	8.59	-1
170	417078	15,16	8	1	8.46	-1
170	417078	15,17	8	0	8.59	-1
176	446531	16,17	0	6	-1	5.91
178	380127	28,29	9	26	-1	2,32
199	468109	15,17	6	0	6.44	-1
199	468109	27,29	6	0	8.09	-1
199	468109	28,29	7	0	8.72	-1
200	447326	15,17	6	0	6.44	-1
219	650195	15,16	6	0	6.34	-1
219	650195	15,17	6	0	6.44	-1
238	470462	15,17	7	0	7.51	-1
242	466697	12,13	6	0	6.2	-1
248	447147	27.29	9	0	12.14	-1
255	447750	12,14	9	0	9,43	-1
255	447750	27,29	9	0	12.14	-1
255	447750	28,29	5	0	6.23	-1
261	560868	27,29	6	0	8.09	-1
276	640356	15.17	10	0	10.73	-1
287	649852	15,17	6	0	6.44	-1
292	446974	28.29	5	0	6.23	-1
307	643924	15.16	6	0	6.34	-1
307	643924	15,17	6	0	6.44	-1
309	452986	15,16	8	ī	8.46	-1
320	449861	16,17	74	26	2.89	-1
320	449861	27,29	1	16	-1	11,86
326	450225	15,17	11	3	3,94	-1
326	450225	27,28	1	24	-1	22.16
326	450225	28,29	24	1	29.9	-1
327	452707	30,31	7	0	7.79	-1
337	452204	15,16	9	1	9.51	-1
337	452204	16,17	1	8	-1	7,88
337	452204	23,24	3	13	-1	4.29
337	452204	27,28	0	14	-1	12,93
337	452204	28,29	14	0	17.44	-1
343	639662	15,16	7	. 0	7.4	-1
343	639662	15,17	7	0	7.51	-1
351	448606	15,17	5	20	-1	3,73
351	448606	27,28	0	7	-i	6.46
374	475184	25,26	6	74	-1	12.78
374	475184	27,28	114	48	2,57	-1
374	475184	27,29	114	70	2.2	-1
374			5		-1	3.42
374 403	475184 555193	30,31 15,16	5	19 0	-1 6.34	3.42 -1

		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
417	3	01,02	5284	2168	2.25	-1
417	3	08,09	1095	631	2.43	-1
417	3	18,20	2079	463	3.84	-1
417	3	19,20	1419	463	2.29	-1
417	3	27,29	993	3994	-1	2.98
417	3	28,29	786	3994	-1	4.08
419	237288	15,16	7	Ō	7.4	-1
419	237288	15,17	7	0	7.51	-1
432	639459	15,17	11	0	11.81	-1
432	639459	16,17	9	0	9.14	-1
467	639480	27,28	8	0	8.66	-1
467	639480	27,29	8	0	10.79	-1
468	644242	15,16	6	0	6.34	-1
468	644242	15,17	6	0	6.44	-1
475	640747	15,17	6	0	6.44	-1
488	640356	15,17	10	0	10.73	-1
490	31112	15,17	6	0	6.44	-1
490	31112	23,24	107	9	12.01	-1
500	23961	15,16	102	44	2.45	-1
500	23961	15,17	102	40	2,74	-1
500	23961	25,26	73	169	-l	2.4
514	645538	23,24	0	6	-1	5,94
517	556	25,26	29	57	-1	2.04
517	556	27,28	34	14	2.63	-1
517	556	28,29	14	43	-1	2.47
517	556	30,31	29	105	-1	3.26
554	446371	15,16	6	0	6.34	-1
558	640221	15,17	8	0	8.59	-l
566	5201	03,04	11	2	5,37	-1
566	5201	15,16	17	4	4.49	-1
566	5201	15,17	17	1	18.25	-1
566	5201	23,24	8	1	8.08	-1
566	5201	27,28	9	0	9.75	-1
566	5201	27,29	9	0	12.14	-1
570	639480	27,28	- 8	0	8.66	-1
570	639480	27,29	8	0 .	10.79	-1
577	649717	16,17	6	0	6.09	-1
587	557401	15,17	10	2	5.37	-1
609	650204	15,17	11	0	11.81	-1
624	645073	15,16	6	0	6.34	-1
624	645073	15,17	6	0	6.44	-1
637	981	18,19	1	9	-1	7.88
637	981	27,28	55	17	3.5	-1
637	981	28,29	17	47	-l	2.22
637	981	30,31	31	224	-1	6.5
661	448450	27,28	0	7	-1	6.46
661	448450	28,29	7	0	8.72	1
662	643804	15,16	6	0	6.34	-1

		PAIR		l l	RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
662	643804	15,17	6	0	6.44	-1
672	607430	15,16	6	0	6.34	-1
678	466697	12,13	6	0	6.2	-1
681	462659	15,16	7	0	7.4	-1
681	462659	15,17	7	0	7.51	-1
683	24730	21,22	0	12	-1	12.2
716	638854	15,16	32	11	3.07	-1
716	638854	15,17	32	0	34.35	-1
716	638854	16,17	11	0	11,17	-1
720	643594	15,16	7	0	7.4	-1
720	643594	15,17	7	0	7.51	-1
729	644442	27,28	0	7	-1	6.46
729	644442	28,29	7	0	8.72	-1
735	447035	16,17	8	1	8.12	-1
743	380550	27,28	14	5	3.03	-1
743	380550	27,29	14	0	18.89	-1
743	380550	28,29	5	0	6.23	-1
744	645538	23,24	0	6	-1	5.94
746	650773	15,17	6	0	6.44	-1
761	650517	15,16	6	0	6.34	-1
761	650517	15,17	6	0	6.44	-1
769	640356	15,17	10	0	10.73	-1
773	450559	16,17	2	11	-1	5.42
797	644242	15,16	6	0	6.34	-1
797	644242	15,17	6	0	6.44	-1
798	644407	30,31	6	0	6.67	-1
814	649170	27,29	6	0	8.09	-1
826	447326	15,17	6	0	6.44	-1
832	400741	23,24	37	14	2.67	-1
832	400741	30,31	19	2	10,57	-1
855	647952	15,16	6	0	6.34	-1
855	647952	15,17	6	0	6.44	-1
855	647952	28,29	8	1	9.97	-1
858	455413	23,24	4	17	-1	4.21
866	645092	15,17	8	0	8.59	-1
873	204	03,04	120	318	-1	2.72
873	204	15,17	35	82	-1	2.18
873	204	27,28	41	14	3.17	-1
873	204	27,29	41	24	2.3	-1
873	204	30.31	287	56	5.7	-1
887	645900	15,17	9	0	9,66	-1
887	645900	16,17	9	0	9.14	-i
887	645900	23,24	7	0	7.07	-1
905	557852	28,29	6	1 0	7.48	-1
918	451709	27,28	2	31	-1	14.31
918	451709	28,29	31	3	12.87	-1
939	448358	27,28	28	15	2.02	-1
945	639480	27,28	8	0	8.66	-1

		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
945	639480	27,29	8	0	10.79	-1
946	1318	01,02	17	42	-1	2.68
946	1318	03,04	42	88	-1	2.15
957	451361	23,24	1	- 8	-1	7.92
958	449891	15,16	8	1	8.46	-1
986	418763	15,16	15	3	5.28	-1
986	418763	15,17	15	2	8.05	-1
986	418763	27,29	11	0	14.84	-1
986	418763	28,29	5	. 0	6.23	-1
987	619635	25,26	7	0	6.76	-1
989	451899	15,17	9	1	9.66	-1
989	451899	27,28	4	21	-1	4.85
991	560860	27,28	0	7	-1	6.46
991	560860	28,29	7	0	8.72	-1
996	452775	27,28	1	9	-1	8.31
1006	649556	28,29	6	0	7.48	-1
1007	644611	15,16	7	0	7.4	-1
1007	644611	15,17	7	0	7.51	-1
1009	470462	15,17	7	0	7.51	-1
1010	645662	23,24	4	28	-1	6.93
1011	649259	15,16	7	0	7.4	-1
1011	649259	15,17	7	0	7.51	-1
1011	649259	27,28	8	0	8.66	-1
1011	649259	27,29	8	0	10.79	-1
1034	532307	16,17	0	7	-1	6.89
1044	414739	15,17	6	19	-1	2.95
1044	414739	16,17	2	19	-1	9.35
1047	238586	30,31	9	0	10.01	-1
1068	467057	23,24	0	7	-1	6.93
1090	549786	23,24	0	8	-1	7.92
1101	649259	15,16	7	0	7.4	-1
1101	649259	15,17	7	0	7.51	-1
1101	649259	27,28	8	0	8.66	-1
1101	649259	27,29	8	0	10.79	-1
1106	468689	15,16	6	0	6.34	-1
1106	468689	15,17	6	0	6.44	-1
1112	556325	16,17	1	8	-1	7.88
1123	645530	15,16	10	2	5.28	-1
1123	645530	15,17	10	0	10.73	-1
1132	452026	15,16	38	14	2.87	-1
1132	452026	27,28	7	26	-l	3.43
1132	452026	28,29	26	2	16.2	-1
1132	452026	30,31	9	2	5.01	-1
1134	612572	15,16	6	0	6.34	-1
1147	14157	28,29	6	0	7.48	-1
1151	454906	27,28	4	18	-l	4.16
1151	454906	28,29	18	2	11.21	-1
1154	62053	25,26	10	1	9.65	-1

		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
1180	645900	15,17	9	0	9.66	-1
1180	645900	16,17	9	0	9.14	-1
1180	645900	23,24	7	0	7.07	-1
1185	463824	15,16	6	0	6.34	-1
1185	463824	15,17	6	0	6.44	-1
1185	463824	27,28	0	- 8	-1	7.39
1185	463824	28,29	8	0	9.97	-1
1193	649617	28,29	5	0	6.23	-1
1208	452738	28,29	5	0	6.23	-1
1234	647232	16,17	9	0	9.14	-1
1234	647232	28,29	6	0	7.48	-1
1237	503122	25,26	8	178	-1	23.05
1237	503122	27,29	12	2	8.09	-1
1237	503122	30,31	22	98	-1	4
1238	515350	15,16	14	0	14.8	-1
1238	515350	15,17	14	3	5.01	-1
1244	648996	15,16	6	0	6.34	-1
1244	648996	15,17	6	0	6.44	-1
1255	416624	27,29	6	0	8.09	-1
1261	449956	12,13	6	0	6.2	-1
1261	449956	16,17	10	1	10.16	-1
1261	449956	28,29	10	0	12.46	-1
1261	449956	30,31	8	1	8.9	-1
1270	380477	15,17	7	0	7.51	-1
1276	645100	15,16	7	0	7.4	-1
1276	645100	15,17	7	0	7.51	-1
1278	554581	28,29	7	0	8.72	-1
1290	650820	16,17	8	0	8.12	-1
1306	646309	16,17	6	0	6.09	-1
1315	502683	15,16	6	0	6.34	-1
1315	502683	28,29	5	0	6.23	-1
1342	463487	15,17	8	1	8.59	-1
1352	446987	15,17	10	0	10.73	-1
1354	640922	27,28	0	7	-1	6.46
1354	640922	28,29	7	0	8.72	-1
1355	561793	30,31	6	0	6.67	-1
1382	649354	15,16	6	0	6.34	-1
1382	649354	15,17	6	0	6.44	-1
1386	507050	27,29	9	0	12.14	-1
1386	507050	28,29	7	0	8.72	-1
1392	649272	16,17	8	0	8.12	-1
1412	453470	15,16	12	_ 1	12.68	-1
1412	453470	15,17	12	1	12.88	-1
1423	419255	15,16	11	0	11.63	-1
1423	419255	15,17	11	1	11.81	-1
1424	648996	15,16	6	0	6.34	-1
1424	648996	15,17	6	0	6.44	-1
1425	451361	23,24	1	8	-1	7.92

		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
1426	15296	27,28	6	0	6.5	-1
1429	643327	28,29	10	_ 0	12.46	-1
1432	452646	28,29	5	0	6,23	-1
1442	651073	15,16	7	_ 0	7.4	-1
1442	651073	15,17	7	0	7.51	-1
1442	651073	27,28	0	9	-1	8.31
1442	651073	28,29	9	2	5.61	-1
1452	213	03,04	17	4	4.15	-1
1452	213	08,09	137	403	-1	2.1
1452	213	21,22	2	14	-1	7.12
1452	213	27,28	4	43	-1	9.93
1452	213	27,29	4	47	-1	8.71
1452	213	30,31	123	35	3.91	-1
1468	268336	15,16	8	1	8.46	-1
1478	646060	15,16	13	3	4.58	-1
1478	646060	15,17	13	0	13.95	-1
1497	639378	27,28	8	0	8.66	-1
1497	639378	27,29	8	0	10.79	-1
1507	446910	25,26	80	346	-1	4.48
1507	446910	27,29	210	38	7.46	-1
1507	446910	28,29	210	38	6.88	-1
1508	447126	15,16	19	5	4.02	-1
1508	447126	16,17	5	20	-1	3,94
1508	447126	28,29	5	0	6.23	-1
1514	479131	23,24	1	13	-1	12.87
1523	284586	15,17	8	0	8.59	-1
1523	284586	27,28	0	17	-1	15.7
1523	284586	28,29	17	0	21.18	-1
1529	449437	15,16	14	3	4.93	-1
1529	449437	16,17	3	12	-1	3.94
1529	449437	27,28	0	8	-1	7.39
1529	449437	28,29	8	0	9.97	-1
1552	449438	27,29	5	0	6.75	-1
1552	449438	28,29	9	0	11.21	-1
1561	448152	27,28	1	17	-1	15.7
1561	448152	28,29	17	2	10,59	-1
1564	479880	23,24	0	7	-1	6.93
1594	449521	27,29	7	i	9.44	-1
1602	644190	16,17	6	0	6.09	-1
1620	639991	15,17	6	0	6.44	-1
1638	594994	28,29	6	0	7.48	-1
1660	376342	15,16	0	7	-1	6.62
1662	734646	15,16	0	14	-1	13,25
1662	734646	16,17	14	0	14.22	-1
1666	460284	23,24	6	0	6.06	-1
1666	460284	27,28	6	0	6.5	-1
1685	558412	30,31	6	0	6.67	-1
1689	640158	16,17	6	0	6.09	-1

	100	PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
1706	209	01,02	9	36	-1	4.34
1706	209	03,04	445	205	2.12	-1
1706	209	15,16	36	188	-1	4.94
1706	209	15,17	36	145	-1	3.75
1706	209	18,19	0	12	-1	10.5
1706	209	19,20	12	1	8.98	-1
1706	209	21,22	19	101	-1	5.4
1706	209	27,28	250	36	7.52	-1
1706	209	27,29	250	69	4.89	-1
1713	423420	30,31	12	0	13.35	-1
1717	448758	23,24	12	27	-1	2.23
1736	470462	15,17	7	0	7.51	-1
1740	729779	15,16	0	8	-1	7.57
1740	729779	16,17	8	0	8.12	-1
1758	649722	15,17	8	0	8.59	-1
1758	649722	16,17	6	0	6.09	-1
1759	562137	23,24	8	0	8.08	-1
1769	27083	16,17	11	0	11.17	-1
1769	27083	25,26	17	56	-1	3.41
1769	27083	28,29	1	14	-1	11.24
1769	27083	30,31	31	99	-1	2.87
1784	498194	27,28	0	7	-1	6.46
1784	498194	28,29	7	0	8.72	-1
1813	734646	15,16	0	14	-1	13.25
1813	734646	16,17	14	0	14.22	-1
1816	730282	15,16	0	7	-1	6.62
1816	730282	16,17	7	0	7.11	-1
1834	640116	15,16	19	4	5.02	-1
1834	640116	15,17	19	0	20.39	-1
1842	465446	25,26	26	11	2.28	-1
1879	27083	16,17	11	0	11.17	-1
1879	27083	25,26	17	56	-1	3.41
1879	27083	28,29	1	14	-1	11.24
1879	27083	30,31	31	99	-1	2.87
1880	478458	27,29	5	0	6.75	-1
1892	734622	23,24	5	17	-1	3.37
1896	381623	15,16	10	2	5.28	-1
1896	381623	15,17	10	0	10.73	-1
1898	446575	27,28	1	10	-1	9.23
1898	446575	28,29	10	1	12.46	-1
1940	649106	15,17	7	0	7.51	-1
1971	727760	27,28	0	- 8	-1	7.39
1971	727760	28,29	8	0	9,97	-1
1972	454087	15,16	14	3	4.93	-1
1972	454087	15,17	14	1	15.03	-1
1972	454087	25,26	4	14	-1	3.63
1978	4584	01,02	1	11	-1	11.93
1978	4584	25,26	6	0	5.79	-1

		PAIR.			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS_	MINUS
2009	734622	23,24	5	17	-1	3.37
2027	48619	30,31	6	0	6.67	-1
2033	646309	16,17	6	0	6.09	-1
2040	447150	23,24	_ O	6	-1	5.94
2041	728273	15,16	0	7	-1	6.62
2041	728273	16,17	7	0	7.11	-1
2048	438663	15,16	4	16	-1	3.78
2048	438663	25,26	97	210	-1	2.24
2048	438663	27,28	48	171	-1	3.29
2048	438663	28,29	171	58	3.67	-1
2048	438663	30,31	13	30	-1	2.07
2064	128773	15,16	4	25	-1	5.91
2064	128773	16,17	25	2	12.69	-1
2064	128773	25,26	79	261	-1	3.42
2064	128773	30,31	13	33	-1	2.28
2074	504513	15,17	6	0	6.44	-1
2082	549801	28,29	8	0	9.97	-1
2083	730484	23,24	1	12	-1	11.88
2100	606076	23,24	2	22	-1	10.89
2122	732712	15,16	0	7	-1	6.62
2122	732712	16,17	7	0	7.11	-1
2132	730059	15.16	0	6	-1	5.68
2132	730059	16,17	6	0	6.09	-1
2137	451184	16.17	14	4	3.55	-1
2140	152	03,04	153	566	-1	3.79
2140	152	08,09	136	395	-1	2.08
2140	152	15.16	59	194	-1	3.11
2140	152	15,17	59	455	-1	7.18
2140	152	16,17	194	455	-1	2.31
2140	152	21,22	232	565	-1	2.48
2140	152	30,31	75	211	-1	2.53
2150	648774	15.17	6	0	6.44	-1
2157	548275	25.26	16	3	5.15	-1
2159	479572	25,26	20	53	-1	2.75
2159	479572	28.29	96	55	2.17	-1
2200	448046	15,17	0	12	-1	11.18
2200	448046	16,17	3	12	-1	3.94
2200	448046	27,28	1	9	-1	8.31
2200	448046	28,29	9	0	11.21	-1
2204	732756	28.29	6	0	7.48	-1
2215	171511	15.16	0	7	-1	6.62
2215	171511	16,17	7	0	7.11	-1
2232	203793	16,17	12	3	4.06	-1
2232	203793	25,26	12	188	-1	16.23
2232	203793	27,28	185	66	3.04	-1
2232	203793	27,29	185	100	2,5	-1
2236	725825	15,16	0	8	-1	7.57
2236	725825	16,17	8	0 1	8.12	-1

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		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
2276	407723	12,13	6	0	6.2	-l
2285	411128	12,13	24	- 8	3.1	-1
2285	411128	13,14	8	29	-l	3.57
2285	411128	18,19	387	158	2.8	-1
2285	411128	19,20	158	634	-l	5.36
2285	411128	25,26	6272	2943	2.06	-1
2285	411128	27,29	4900	2229	2.97	-l
2285	411128	28,29	4750	2229	2.65	-l
2289	726173	15,16	0	6	-1	5.68
2289	726173	16,17	6	0	6.09	-1
2322	732712	15,16	0	7	-1	6.62
2322	732712	16,17	7	0	7.11	-1
2337	4255	08,09	28	7	5.59	-1
2338	554080	16,17	_1	9	-1	8.86
2358	4244	28,29	- 8	28	7	2.81
2376	737087	28,29	5	0	6.23	-1
2378	735871	23,24	0	13	-1	12.87
2388	735292	15,16	0	9	-1	8.51
2388	735292	16,17	9	0	9.14	-1
2400	732223	27,29	5	0	6.75	-1
2401	595506	15,16	0	6	-1	5.68
2405	728884	15,16	0	8	-1	7.57
2405	728884	16,17	8	0	8.12	-1
2417	453508	25,26	27	9	2.9	-1
2417	453508	27,28	9	1	9.75	-l
2442	620462	23,24	2	19	-1	9.4
2445	735028	15,16	0	6	-1	5.68
2445	735028	16,17	_6	0	6.09	-l
2449	447075	27,28	3	37	-1	11.39
2449	447075	28,29	37	14	3.29	-l
2483	645139	16,17	9	0	9.14	-1
2493	401368	27,29	48	32	2.02	-l
2514	451764	16,17	16	5	3,25	-1
2519	105056	15,16	1	10	-l	9.46
2529	560868	27,29	6	0	8.09	-1
2532	1030	21,22	13	3	4.26	-1
2532	1030	27,29	21	12	2.36	-l
2532	1030	30,31	14	. 5	3.11	-1
2545	455581	27,28	118	49	2.61	-1
2545	455581	27,29	118	29	5.49	-1
2545	455581	28,29	49	29	2.11	-1
2573	447550	23,24	6	0	6.06	-1
2573	447550	25,26	48	197	-1	4.25
2573	447550	30,31	6	20	-1	3
2591	447003	28,29	6	0 .	7.48	-1
2597	649852	15,17	6	0	6.44	-1
2598	650297	15,17	6	0	6.44	-1
2609	402516	30,31	6	0	6.67	-1

CEO ID VO	CI LIOTER	PAIR	er come :	GT 03775	RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
2629	405016	16,17	15	4	3.81	-1
2655	734031	27,28	0	7	-1	6.46
2655	734031	28,29	7	0	8.72	-1
2663	736595	15,16	0	6	-1	5.68
2663	736595	16,17	6	0	6.09	-1
2669	734209	15,16	0	9	-1	8.51
2669	734209	16,17	9	0	9.14	-1
2673	561632	27,29	5	0	6.75	-1
2681	530883	15,16	0	7	-1	6.62
2682	729173	15,16	0	8	-1	7.57
2682	729173	16,17	8	0	8.12	-1
2685	726786	12,13	6	0	6.2	-1
2685	726786	15,16	0	10	-1	9.46
2685	726786	16,17	10	0	10.16	-1
2685	726786	30,31	6	0	6.67	-1
2687	448770	15,17	6	0	6.44	-1
2712	726449	27,28	16	1	17.33	-1
2712	726449	27,29	16	0	21.59	-1
2718	446752	12,14	1	12	-1	11.45
2750	289328	12,13	6	0	6.2	-1
2750	289328	30,31	8	·1	8.9	-1
2772	81	15,16	30	79	-1	2.49
2772	81	15,17	30	114	-1	3.54
2772	81	18,19	0	46	-1	40.25
2772	81	19,20	46	4	8.6	-1
2772	81	27,29	49	175	-1	2.65
2772	81	30,31	52	116	-1	2.01
2775	220107	15,17	9	25	-1	2.59
2775	220107	28,29	9	39	-1	3.48
2780	402024	12,13	2	12	-1	5,81
2780	402024	12,14	2	11	-1	5.25
2782	451888	27,29	79	26	4.1	-1
2782	451888	28,29	70	26	3.35	-1
2791	731467	15,16	0	6	-1	5.68
2791	731467	16,17	6	0	6.09	-1
2801	724781	28,29	10	2	6,23	-1
2807	553850	28,29	5	0	6.23	-1
2849	555103	15,17	0	7	-1	6.52
2849	555103	23,24	0	6	-1	5.94
2861	735477	15,16	0	7	-1	6.62
2861	735477	16,17	7	0	7.11	-1
2878	736014	15,16	0	6	-1	5,68
2878	736014	16,17	6	0	6.09	-1
2879	42	08,09	190	121	2.19	-1
2879	42	27,28	792	286	3	-1
2879	42	28,29	286	1257	-1	3,53
2882	42	08,09	190	121	2.19	-1
2882	42	27,28	792	286	3	-1

SEC ID NO   CLUSTER   AB   CLONES A   CLONES B   PLUS   MINUS							
2882         42         28.29         286         1257         -1         3.53           2883         736014         15,16         0         6         -1         5.68           2883         736014         16,17         6         0         6.09         -1           2885         4470         03,04         8         0         7.81         -1           2885         4470         15,16         10         24         -1         227           2885         4470         15,17         10         0         10,73         -1           2885         4470         15,17         10         0         24,37         -1           2885         4470         15,17         0         9         -1         8.38           2897         554703         15,17         0         9         -1         8.38           2897         554703         15,17         10         1         10,73         -1           2898         546642         15,17         10         1         10,73         -1           2898         546642         15,17         22         1         22,24         -1           2910<							
2883   736014   15,16   0   6   1   5,68							
2883         736014         16,17         6         0         6,09         -1           2885         4470         03,04         8         0         7,81         -1           2885         4470         15,16         10         24         -1         2,27           2885         4470         15,17         10         0         10,73         -1           2885         4470         15,17         10         0         24,37         -1           2885         4470         25,26         7         0         6,76         -1           2887         554703         15,17         0         9         -1         8,36           2898         546642         15,17         10         1         10,73         -1           2898         546642         15,17         10         1         10,73         -1           2910         732300         15,16         0         7         -1         6,62           2910         732300         15,16         0         6         -1         5,68           2938         892         15,16         0         6         -1         5,68           2938							
2885         4470         03,04         8         0         7,81         -1           2885         4470         15,16         10         24         -1         2,27           2885         4470         15,17         10         0         10,73         -1           2885         4470         16,17         24         0         24,37         -1           2885         4470         15,17         0         9         -1         8,38           2897         554703         15,17         0         9         -1         8,38           2898         546642         15,17         10         1         10,73         -1           2898         546642         16,17         22         1         22,34         -1           2910         732,300         15,16         0         7         -1         6,62           2910         732,300         15,16         0         6         -1         5,68           2938         892         27,28         6         0         6,9         -1           2938         892         27,28         6         0         6,9         -1           2938							5.68
2885         4470         15,16         10         24         -1         2,27           2885         4470         15,17         10         0         10,73         -1           2885         4470         15,17         24         0         24,37         -1           2885         4470         25,26         7         0         6,76         -1           2897         554703         16,17         1         9         -1         8,38           2898         546642         15,17         10         1         10,73         -1           2910         732300         15,16         0         7         -1         6,62           2910         732300         15,16         0         7         -1         6,62           2910         732300         15,16         0         6         -1         5,68           2938         892         15,16         0         6         -1         5,68           2938         892         15,16         0         6         -1         5,68           2938         892         15,16         0         6         -1         5,68           2938							
2885         4470         15,17         10         0         10,73         -1           2885         4470         16,17         24         0         24,37         -1           2885         4470         25,26         7         0         6,76         -1           2897         554703         15,17         0         9         -1         8,38           2898         546642         15,17         10         1         10,73         -1           2898         546642         15,17         10         1         10,73         -1           2910         732300         15,16         0         7         -1         6,62           2910         732300         15,16         0         7         -1         6,62           2938         892         15,16         0         6         -1         5,68           2938         892         27,29         6         0         8,09         -1           2938         892         27,29         6         0         8,09         -1           2941         546642         15,17         2         1         10,73         -1           2941							
2885         4470         16,17         24         0         24,37         -1           2885         4470         16,17         24         0         24,37         -1           2887         54703         15,17         0         9         -1         8,38           2897         554703         16,17         1         9         -1         8,38           2898         546642         16,17         22         1         10,73         -1           2898         546642         16,17         22         1         22,34         -1           2910         732300         15,16         0         7         -1         6,62           2910         732300         15,16         0         6         -1         5,68           2938         892         15,16         0         6         -1         5,68           2938         892         27,28         6         0         6.5         -1           2938         892         27,29         6         0         8,09         -1           2941         546642         15,17         10         1         10,73         -1           2949							2,27
2885         4470         25,26         7         0         6,76         -1           2897         554703         15,17         0         9         -1         8,38           2898         554703         16,17         1         9         -1         8,36           2898         546642         15,17         10         1         10,73         -1           2898         546642         15,17         10         1         10,73         -1           2910         732300         15,16         0         7         -1         6,62           2910         732300         15,16         0         6         -1         5,68           2938         892         15,16         0         6         -1         5,68           2938         892         27,28         6         0         6.5         -1           2938         892         27,29         6         0         8.09         -1           2941         546642         15,17         10         1         10,73         -1           2941         546642         15,17         2         1         22,24         -1           2941							
2897         554703         15,17         0         9         -1         8,38           2897         554703         16,17         1         9         -1         8,36           2898         5546642         15,17         10         1         10,73         -1           2898         546642         16,17         22         1         22,34         -1           2910         732300         15,16         0         7         -1         6,62           2910         732300         16,17         7         0         7,11         -1           2938         892         15,16         0         6         -1         5,68           2938         892         15,16         0         6         -1         5,68           2938         892         27,28         6         0         6,0         -1         5,68           2938         892         27,29         6         0         8,09         -1         1         10,73         -1           2941         546642         15,17         10         1         10,73         -1         22,34         -1         5,08         -1         2949         734832 <td></td> <td></td> <td></td> <td></td> <td></td> <td>24.37</td> <td>-1</td>						24.37	-1
2897         554703         16,17         1         9         -1         8.86           2898         546642         15,17         10         1         10,73         -1           2898         546642         16,17         22         1         22,34         -1           2910         732300         15,16         0         7         -1         6.62           2910         732300         15,16         0         6         -1         5.68           2938         892         15,16         0         6         -1         5.68           2938         892         15,16         0         6         -1         5.68           2938         892         27,29         6         0         8.09         -1           2941         546642         15,17         10         1         10,73         -1           2941         546642         15,17         10         1         10,73         -1           2949         734382         15,16         0         6         -1         5.68           2949         734382         15,16         0         6         -1         5.68           2949 <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>							
2898         546642         15,17         10         1         10,73         -1           2898         346642         16,17         22         1         22,34         -1           2910         732300         15,16         0         7         -1         6,62           2910         732300         16,17         7         0         7,11         -1           2938         892         15,16         0         6         -1         5,68           2938         892         27,28         6         0         6,5         -1           2938         892         27,29         6         0         8,09         -1           2941         546642         15,17         10         1         10,73         -1           2941         546642         16,17         22         1         22,34         -1           2949         734382         15,16         0         6         -1         5,68           2949         734382         15,17         0         6         -1         5,68           2949         734382         15,17         2         25         -1         11,69           2983 <td></td> <td></td> <td></td> <td>0</td> <td></td> <td>-1</td> <td>8.38</td>				0		-1	8.38
2898   546642   16,17   22   1   22,34   -1							8.86
2910							-1
2910						22.34	
2938         892         15,16         0         6         -1         5.68           2938         892         27,28         6         0         6.5         -1           2938         892         27,29         6         0         8.09         -1           2941         546642         15,17         10         1         10,73         -1           2949         734382         15,16         0         6         -1         5.68           2949         734382         16,17         6         0         6.09         -1           2983         448110         16,17         1         25         -1         11,65           2983         448110         16,17         1         25         -1         24,62           2999         1015         08,09         10         31         -1         2,22           2999         1015         08,09         10         31         -1         2,22           2999         1015         31,4         48         28         2,32         -1           2999         1015         33,1         31         6         5,75         -1           3006		10211					6.62
2938         892         27,28         6         0         6.5         -1           2938         892         27,29         6         0         8.09         -1           2941         546642         15,17         10         1         10.73         -1           2941         546642         16,17         22         1         22,234         -1           2949         734582         15,16         0         6         -1         5.68           2949         734582         16,17         6         0         6.09         -1           2983         448110         15,17         2         25         -1         11.65           2993         448110         16,17         1         25         -1         24.62           2994         450027         23,24         0         6         -1         5.94           2999         1015         08,09         10         31         -1         2.22           2999         1015         13,14         84         28         3.05         -1           3006         402070         27,28         1         9         -1         3.31           3006<							-1
2938   892   27,29   6   0   8.09   -1				0		-1	5.68
2941   \$46642   15,17   10   1   10,73   -1   2941   \$46642   16,17   22   1   22,34   -1   2949   734382   15,16   0   6   -1   5,68   2949   734382   15,16   0   6   -1   5,68   2949   734382   15,17   6   0   6,09   -1   2983   448110   15,17   2   25   -1   11,65   2983   448110   15,17   1   25   -1   24,62   2999   450027   23,24   0   6   -1   5,94   2999   1015   08,09   10   31   -1   2.32   2999   1015   12,14   62   28   2.32   -1   2999   1015   13,14   84   28   3.05   -1   2999   1015   30,31   31   6   5,75   -1   3006   402070   27,28   1   9   -1   3.31   3006   402070   28,29   9   0   11,21   -1   3015   446964   15,17   1   12   -1   11,18   3020   414739   16,17   2   19   -1   2.95   3020   414739   16,17   2   19   -1   9,35   3038   450410   16,17   0   7   -1   6,89   3093   53318   15,16   12   3   4,23   -1   3057   14573   30,31   11   3   4,08   -1   3093   503452   16,17   1   8   -1   7,38   3093   53318   15,17   1   8   -1   7,38   3093   53318   15,17   1   8   -1   7,38   3113   551289   23,24   0   7   -1   6,93   3126   447667   27,28   22   10   2,38   -1   3126   447667   27,28   22   3   9,98   -1   3129   551617   15,16   9   0   9,51   -1   3129   551617   15,16   9   0   9,51   -1   3129   551617   15,16   9   0   9,51   -1   3129   551617   15,16   9   0   9,51   -1   3129   551617   15,16   9   0   9,51   -1   3125   545117   15,17   0   6   -1   5,91			27,28		0	6.5	-1
2941   546642   16,17   22   1   22,34   1   2949   734582   15,16   0   6   -1   5,68   2949   734582   15,16   0   6   -1   5,68   2949   734582   15,17   6   0   6,09   -1   2983   448110   15,17   2   25   -1   11,65   2984   450027   23,24   0   6   -1   5,94   2999   450027   23,24   0   6   -1   5,94   2999   1015   68,09   10   31   -1   2,22   2999   1015   12,14   62   28   2,32   -1   2999   1015   13,14   84   28   3,05   -1   2999   1015   30,31   31   6   5,75   -1   3006   402070   27,28   1   9   -1   8,31   3006   402070   23,29   9   0   11,21   -1   3020   414739   15,17   6   19   -1   2,95   3020   414739   15,17   6   19   -1   2,95   3020   414739   15,17   6   19   -1   2,95   3020   414739   15,17   1   12   -1   30303   430410   16,17   0   7   -1   6,89   3044   553316   15,16   12   3   4,23   -1   3093   503452   16,17   1   3   -1   7,88   3099   533158   16,17   1   8   -1   7,88   3113   554581   28,29   7   0   8,72   -1   3126   447667   27,28   22   10   2,38   -1   3126   447667   27,28   22   3   9,89   -1   3126   447667   27,28   22   10   2,38   -1   3129   551617   15,16   9   0   9,51   -1   3129   551617   15,16   9   0   9,51   -1   3129   551617   15,16   9   0   9,51   -1   3145   551117   16,17   0   6   -1   5,91		892	27,29			8.09	-1
2949   734582   15,16   0   6   -1   5,68	2941	546642	15,17	10	1	10.73	-1
2949         734582         16,17         6         0         6,09         -1           2983         448110         15,17         2         25         -1         11.65           2983         448110         15,17         2         25         -1         11.65           2983         448110         16,17         1         25         -1         24.62           2994         450027         23,24         0         6         -1         5.94           2999         1015         08,09         10         31         -1         2.22           2999         1015         13,14         84         28         3.05         -1           3006         402070         27,28         1         9         -1         8,31           3006         402070         28,29         9         0         11,21         -1           3015         446964         15,17         1         12         -1         11,18           3020         414739         15,17         6         19         -1         2,95           3038         450410         16,17         2         19         -1         9,35	2941	546642	16,17	22	1	22,34	-1
2983	2949	734582	15,16	0	6	-1	5.68
2983         448110         16,17         1         25         -1         24,62           2994         450027         23,24         0         6         -1         5,94           2999         1015         68,09         10         31         -1         2,22           2999         1015         13,14         84         28         2,32         -1           2999         1015         33,14         84         28         3,05         -1           2999         1015         30,31         31         6         5,75         -1           3006         402070         27,28         1         9         -1         8,31           3006         402070         28,29         9         0         11,21         -1         -1           3015         446964         15,17         1         12         -1         -11,18           3020         414739         16,17         2         19         -1         9,35           3038         450410         16,17         0         7         -1         6,89           3044         533316         15,16         12         3         4,23         -1	2949	734582	16,17	6	0	6.09	-1
2994         450027         23,24         0         6         -1         5,94           2999         1015         68,09         10         31         -1         2,22           2999         1015         12,14         62         28         2,32         -1           2999         1015         13,14         84         28         3,05         -1           2999         1015         30,31         31         6         5,75         -1           3006         402070         27,28         1         9         -1         8,31           3006         402070         23,29         9         0         11,21         -1           3020         444739         15,17         1         12         -1         11,18           3020         444739         15,17         6         19         -1         2,95           3038         4850410         16,17         2         19         -1         9,35           3044         553316         15,16         12         3         4,23         -1           3093         503452         16,17         1         8         -1         7,88           3	2983	448110	15,17	2	25	-1	11.65
2999         1015         08,09         10         31         -1         2.22           2999         1015         12,14         62         28         2.32         -1           2999         1015         13,14         84         28         3.05         -1           2999         1015         30,31         31         6         5.75         -1           3006         402070         22,28         1         9         -1         8.31           3006         402070         28,29         9         0         11,21         -1         11,18           3015         446964         15,17         1         12         -1         -11,18           3020         44739         15,17         2         19         -1         9.35           3020         44739         16,17         2         19         -1         9.35           3044         553316         15,16         12         3         4,23         -1           3053         34542         16,17         0         7         -1         6,89           3093         503452         16,17         1         8         -1         7,88	2983	448110	16,17	1	25	-1	24.62
2999         1015         12,14         62         28         2,32         -1           2999         1015         13,14         84         28         3,05         -1           2999         1015         30,31         31         6         5,75         -1           3006         402070         27,28         1         9         -1         8,31           3006         402070         28,29         9         0         11,21         -1           3015         446964         15,17         1         12         -1         11,18           3020         414739         15,17         6         19         -1         2,95           3030         444739         15,17         6         19         -1         2,95           3020         414739         15,17         0         7         -1         6,89           3044         53316         15,16         12         3         4,23         -1           3047         14573         30,31         11         3         4,08         -1           3093         593452         16,17         1         8         -1         7,88           309	2994	450027	23,24	0	6	-1	5.94
2999         1015         13,14         84         28         3,05         -1           2999         1015         30,31         31         6         5,75         -1           3006         402070         22,28         1         9         -1         8,31           3006         402070         28,29         9         0         11,21         -1           3015         446964         15,17         1         12         -1         11,18           3020         414739         15,17         6         19         -1         2,95           3038         430410         16,17         0         7         -1         6,89           3044         553316         15,16         12         3         4,23         -1           3057         14573         30,31         11         3         4,08         -1           3093         503452         16,17         1         8         -1         7,88           3099         553185         16,17         1         8         -1         7,88           3099         553188         16,17         1         8         -1         7,88           312	2999	1015	08,09	10	31	-1	2.22
2999         1015         30,31         31         6         5,75         -1           3006         402070         27,28         1         9         -1         8,31           3006         402070         28,29         9         0         11,21         -1           3015         446964         15,17         1         12         -1         11,18           3020         414739         15,17         6         19         -1         2,95           3020         414739         16,17         2         19         -1         9,35           3038         450410         16,17         0         7         -1         6,89           3044         553316         15,16         12         3         4,23         -1           3053         503452         16,17         1         8         -1         7,88           3093         503452         16,17         1         8         -1         7,88           3099         553138         16,17         1         8         -1         7,88           3113         554289         23,24         0         7         -1         6,93           31	2999	1015	12,14	62	28	2.32	-1
3006   402070   27,28   1   9   -1   8.31   3006   402070   28,29   9   0   11,21   -1   3015   446964   15,17   1   12   -1   11,18   3020   414739   15,17   6   19   -1   2.95   3020   414739   16,17   2   19   -1   9.35   3020   414739   16,17   2   19   -1   9.35   3038   450410   16,17   0   7   -1   6.89   3044   553316   15,16   12   3   4.23   -1   3057   14573   30,31   11   3   4.03   -1   3093   503452   16,17   1   8   -1   7.88   3099   533158   16,17   1   8   -1   7.88   3099   533158   16,17   1   8   -1   7.88   3113   551289   23,24   0   7   -1   6.93   3126   447667   27,28   22   10   2.38   -1   3126   447667   27,28   22   10   2.38   -1   3126   447667   27,28   22   10   2.38   -1   3126   447667   27,29   22   3   9.89   -1   3126   447667   27,29   22   3   9.89   -1   3126   447667   27,29   22   3   9.89   -1   3126   447667   27,29   22   3   9.89   -1   3126   447667   27,29   20   3   9.51   -1   5129   551617   15,16   9   0   9.51   -1   6.89   3145   551117   16,17   0   6   -1   5.91	2999	1015	13,14	84	28	3.05	-1
3006   402070   28,29   9   0   11,21   -1	2999	1015	30,31	31	6	5.75	-1
3015		402070	27,28			7	8.31
3020	3006	402070	28,29	9	0	11.21	-1
3020		446964	15,17	1	12	-1	11.18
3038			15,17		19		2.95
3044   553316   15,16   12   3   4,23   -1	3020	414739	16,17			-1	9.35
3057   14573   30,31   11   3   4.08   -1   7.88   3093   503452   16,17   1   8   -1   7.88   3099   553158   16,17   1   8   -1   7.88   3113   551289   23,24   0   7   -1   6.93   3121   554381   28,29   7   0   8.72   -1   3126   447667   16,17   0   11   -1   10,83   3126   447667   27,28   22   10   2.38   -1   3126   447667   27,29   22   3   9.89   -1   3126   447667   27,29   10   3   4.15   -1   3129   551617   15,16   9   0   9.51   -1   3129   551617   15,16   9   0   9.51   -1   3129   551617   16,17   0   7   -1   6.89   3145   551117   16,17   0   6   -1   5.91							
3093   503452   16,17   1   8   -1   7.88   3099   533158   16,17   1   8   -1   7.88   3099   533158   16,17   1   8   -1   7.88   3113   551289   23,24   0   7   -1   6,93   3121   554581   28,29   7   0   8,72   -1   3126   447667   16,17   0   11   -1   10,83   3126   447667   27,28   22   10   2,38   -1   3126   447667   27,29   22   3   9,89   -1   3126   447667   27,29   10   3   4,15   -1   3129   551617   15,16   9   0   9,51   -1   3129   551617   16,17   0   7   -1   6,89   3145   551117   16,17   0   6   -1   5,91		553316	15,16		3	4.23	-1
3099   553158   16,17   1   8   -1   7.88   3113   551289   23,24   0   7   -1   6,93   3121   554581   28,29   7   0   8.72   -1   3126   447667   16,17   0   11   -1   10.83   3126   447667   27,28   22   10   2.38   -1   3126   447667   27,29   22   3   9.89   -1   3126   447667   28,29   10   3   4.15   -1   3129   551617   15,16   9   0   9.51   -1   3129   551617   16,17   0   7   -1   6.89   3145   551117   16,17   0   6   -1   5.91	3057	14573	30,31	11	3	4.08	-1
3113         551289         23,24         0         7         -1         6,93           3121         554381         28,29         7         0         8,72         -1           3126         447667         16,17         0         11         -1         10,83           3126         447667         27,28         22         10         2,38         -1           3126         447667         27,29         22         3         9,89         -1           3126         447667         28,29         10         3         4,15         -1           3129         551617         15,16         9         0         9,51         -1           3129         551617         16,17         0         7         -1         6,89           3145         551117         16,17         0         6         -1         5,91	3093	503452	16,17	1	8	-1	7.88
3121   554381   28,29   7   0   8,72   -1	3099	553158	16,17	1	8	-1	7.88
3126         447667         16,17         0         11         -1         10,83           3126         447667         27,28         22         10         2,38         -1           3126         447667         27,29         22         3         9,89         -1           3126         447667         28,29         10         3         4,15         -1           3129         551617         15,16         9         0         9,51         -1           3129         551617         16,17         0         7         -1         6,89           3145         551117         16,17         0         6         -1         5,91	3113	551289	23,24		7	-1	6.93
3126         447667         27,28         22         10         2.38         -1           3126         447667         27,29         22         3         9,89         -1           3126         447667         28,29         10         3         4,15         -1           3129         551617         15,16         9         0         9,51         -1           3129         551617         16,17         0         7         -1         6,89           3145         551117         16,17         0         6         -1         5,91	3121	554581	28,29	7	0	8.72	-1
3126         447667         27,29         22         3         9,89         -1           3126         447667         28,29         10         3         4,15         -1           3129         551617         15,16         9         0         9,51         -1           3129         551617         16,17         0         7         -1         6,89           3145         551117         16,17         0         6         -1         5,91	3126	447667	16,17	0	11	-1	10.83
3126   447667   28,29   10   3   4.15   -1	3126	447667	27,28	22	10	2.38	-1
3129         551617         15,16         9         0         9,51         -1           3129         551617         16,17         0         7         -1         6.89           3145         551117         16,17         0         6         -1         5.91	3126	447667	27,29	22	3	9.89	-1
3129 551617 16,17 0 7 -1 6.89 3145 551117 16,17 0 6 -1 5.91	3126	447667	28,29	10	3	4.15	-1
3145 551117 16,17 0 6 -1 5.91	3129	551617	15,16	9	0	9.51	-1
	3129	551617	16,17	0	7	-1	6.89
3150 1093 27,28 43 20 2.33 -1	3145	551117		0	6		5.91
	3150	1093	27,28	43	20	2.33	-1

		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
3150	1093	30,31	5	18	-1	3.24
3151	2284	23,24	6	17	-1	2.8
3151	2284	25,26	17	4	4.1	-1
3151	2284	30,31	14	32	-1	2.05
3168	275	15,17	3	15	-1	4.66
3168	275	18,19	0	12	-1	10.5
3172	551718	15,17	0	8	-1	7.45
3192	143346	25,26	44	435	-1	10,24
3211	551553	28,29	7	0	8.72	-1
3248	556216	15,17	0	7	-1	6.52
3248	556216	16,17	0	7	-1	6.89
3291	446814	25,26	10	27	-1	2.8
3291	446814	27,29	8	1	10.79	-1
3301	550694	15,16	21	1	.22.2	-1
3301	550694	15,17	21	7	3.22	-1
3307	450963	28,29	6	0	7.48	-1
3328	554764	25,26	32	0	30.89	-1
3339	550855	16,17	1	10	-1	9.85
3355	452506	15,16	8	1	8.46	-1
3355	452506	28,29	6	0	7.48	-1
3359	408130	30,31	10	0	11.12	-1
3363	549964	23,24	8	1	8.08	-1
3372	380127	28,29	9	26	-1	2.32
3378	549320	15,17	0	7	-1	6.52
3382	558103	27,28	1	11	-1	10.16
3382	558103	28,29	11	0	13.7	-1
3382	558103	30,31	6	0	6.67	-1
3383	548864	30,31	6	0	6,67	-1
3391	10397	03,04	10	0	9.76	-1
3391	10397	27,28	6	0	6.5	-1
3391	10397	27,29	6	0	8.09	-1
3393	561892	15,17	0	7	-1	6.52
3398	548965	16,17	4	14	-1	3.45
3435	549829	16,17	0	6	-1	5.91
3452	257547	23,24	29	8	3.66	-1
3452	257547	27,28	0	8	-1	7.39
3452	257547	28,29	8	0	9.97	-1
3457	450255	16,17	0	6	-1	5.91
3469	448663	28,29	5	0	6.23	-1
3474	559575	27,28	1	9	-1	8.31
3474	559575	28,29	9	0	11.21	-1
3485	4244	28,29	8	28	-1	2.81
3499	349	01,02	78	160	-1	2.22
3499	349	03,04	91	3	29.59	-1
3499	349	12,13	33	13	2.62	-l
3499	349	13,14	13	28	-1	2.12
3499	349	16,17	15	35	-1	2.3
3499	349	27,28	12	29	-1	2.23

		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
3512	498194	27,28	0	7	-1	6.46
3512	498194	28,29	7	0	8.72	-1
3518	453667	23,24	3	12	-1	3.96
3538	275	15,17	3	15	-1	4.66
3538	275	18,19	0	12	-1	10,5
3559	447429	27,28	0	8	-1	7.39
3559	447429	28,29	8	0	9.97	-1
3563	449956	12,13	6	0	6.2	-1
3563	449956	16,17	10	1	10.16	-1
3563	449956	28,29	10	0	12.46	-1
3563	449956	30,31	8	1	8.9	-1
3573	134392	23,24	20	2	10.1	-1
3573	134392	25,26	0	6	-1	6.22
3583	551380	16,17	4	14	-1	3.45
3600	450242	30,31	7	0	7.79	-1
3603	451361	23,24	1	8	-1	7.92
3608	402916	23,24	8	0	8.08	-1
3608	402916	27,28	22	5	4.76	-1
3608	402916	27,29	22	4	7.42	-1
3613	555502	16,17	0	9	-1	8.86
3619	461	15,16	14	39	-1	2.64
3619	461	15,17	14	34	-1	2.26
3619	461	18,19	1	10	-1	8.75
3619	461	19,20	10	0	7.48	-1
3619	461	27,29	41	124	-1	2.24
3619	461	28,29	37	124	-1	2.69
3619	461	30,31	21	119	-1	5.09
3625	450867	15,16	7	0	7.4	-1
3627	502683	15,16	6	0	6.34	-1
3627	502683	28,29	5	0	6.23	-1
3637	447405	15,17	27	83	-1	2.86
3637	447405	18,20	2	11	-1	6.43
3637	447405	19,20	0	11	-1	14.71
3637	447405	27,29	163	60	3.66	-1
3637	447405	30,31	14	36	-1	2.31
3641	560984	16,17	0	6	-1	5.91
3650	554604	15,17	0	7	-1	6.52
3662	447858	15,16	12	3	4.23	-1
3721	555830	15,16	16	2	8.46	-1
3721	555830	15,17	16	2	8.59	-1
3731	562550	16,17	0	6	-1	5.91
3736	873	15,17	6	21	-1	3.26
3736	873	27,29	15	50	-1	2.47
3736	873	28,29	17	50	-1	2,36
3736	873	30,31	14	32	-1	2.05
3744	126	15,16	103	260	-1	2.39
3744	126	15,17	103	571	I-	5.16
3744	126	16,17	260	571	-l	2.16

		PATR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
3744	126	18,19	4	37	-1	8.09
3744	126	18,20	4	13	-1	3,8
3744	126	19,20	37	13	2.13	-1
3744	126	27,28	638	323	2.14	-1
3762	555103	15,17	0	7	-1	6,52
3762	555103	23,24	0	6	-1	5.94
3774	531145	28,29	6	0	7.48	-1
3782	129715	08,09	3	27	-1	6.44
3807	554764	25,26	32	0	30.89	-1
3810	556561	15,17	0	10	-1	9.32
3810	556561	16,17	1	10	-1	9.85
3823	553787	28,29	10	0	12.46	-1
3848	556759	12,13	8	1	8.26	-1
3848	556759	15,16	11	2	5.81	-1
3848	556759	16,17	2	18	-1	8.86
3848	556759	27,29	21	1	28.33	-1
3848	556759	28,29	18	1	22.43	-1
3848	556759	30,31	10	1	11.12	-1
3857	503452	16,17	1	8	-1	7.88
3861	2284	23,24	6	17	-1	2.8
3861	2284	25,26	17	4	4.1	-1
3861	2284	30,31	14	32	-1	2.05
3864	560984	16,17	0	6	-1	5.91
3873	551444	23,24	6	0	6.06	-1
3881	548858	16,17	0	11	-1	10.83
3885	411113	16,17	0	6	-1	5.91
3894	594994	28,29	6	0	7.48	-l
3904	562550	16,17	0	6	-1	5.91
3922	460445	16,17	0	6	-1	5.91
3925	931	01,02	23	59	-1	2.78
3925	931	18,19	0	9	-1	7.88
3925	931	30,31	70	195	-1	2.5
3940	388688	27,29	8	2	5.4	-1
3940	388688	28,29	17	2	10.59	-1
4009	380127	28,29	9	26	-1	2.32
4016	378459	23,24	1	9	-1	8.91
4017	550571	16,17	0	6	-1	5.91
4036	607430	15,16	6	0	6.34	-1
4039	558098	15,16	6	0	6.34	-1
4039	558098	30,31	9	1	10.01	-1
4041	557928	16,17	0	6	-1	5.91
4079	557928	16,17	0	6	-1	5.91
4100	450242	30,31	7	0	7.79	-1
4106	448450	27,28	0	7	-1	6.46
4106	448450	28,29	7	0	8.72	-1
4111	560538	16,17	0	- 8	-1	7.88
4115	420892	13,14	0	6	-1	5.91
4138	450883	28,29	5	0	6,23	-1

		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
4139	638971	15,17	8	0	8.59	-1
4149	643804	15,16	6	0	6.34	-1
4149	643804	15,17	6	0	6.44	-1
4150	97909	27,28	31	12	2.8	-1
4150	97909	28,29	12	35	-1	2,34
4150	97909	30,31	18	54	-1	2.7
4157	603857	27,29	17	3	7.64	-1
4157	603857	28,29	31	3	12.87	-1
4162	641683	15,16	8	0	8,46	-1
4162	641683	15,17	8	0	8.59	-1
4176	640889	15,17	6	0	6.44	-1
4178	643594	15,16	7	0	7.4	-1
4178	643594	15,17	7	0	7.51	-1
4195	641728	28,29	7	1	8.72	-1
4203	359500	12,13	14	4	3.61	-1
4203	359500	27,28	18	4	4.87	-1
4203	359500	28,29	4	17	-1	3.41
4203	359500	30,31	8	22	-1	2.47
4204	649558	15,16	12	3	4.23	-1
4204	649558	15,17	12	0	12.88	-1
4231	451401	15,17	9	1	9.66	-1
4234	417259	15,17	4	14	-1	3.26
4234	417259	16,17	1	14	-1	13.79
4240	453006	27,28	0	8	-1	7.39
4240	453006	28,29	8	0	9.97	-1
4247	498194	27,28	0	7	-1	6.46
4247	498194	28,29	7	0	8.72	-1
4250	556326	16,17	0	8	-1	7.88
4257	649106	15,17	7	0	7.51	-1
4264	284586	15,17	8	0	8.59	-1
4264	284586	27,28	0	17	-1	15.7
4264	284586	28,29	17	0	21.18	-1
4265	642535	27,28	0	8	-1	7.39
4265	642535	28,29	8	0	9.97	-1
4270	801	30,31	25	7	3.97	-1
4271	449956	12,13	6	0	6.2	1
4271	449956	16,17	10	1	10.16	-1
4271	449956	28,29	10	0	12.46	-1
4271	449956	30,31	8	1	8.9	-1
4283	455113	27,29	11	3	4.95	-1
4283	455113	28,29	13	3	5.4	-1
4285	1558	03,04	25	97	-1	3.98
4287	546705	16,17	0	6	-1	5.91
4293	452212	15,16	6	0	6.34	-1
4298	37862	25,26	14	0	13.51	-1
4299	644609	15,17	7	0	7.51	-1
4311	553877	16,17	1	8	-1	7.88
4319	550562	27,28	1	9	-1	8.31

		PAIR			RATIO	RATIO
SEO ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
4319	550562	28.29	9	1	11.21	-1
4320	648320	15.16	6	ô	6.34	-1
4320	648320	15,17	6	0	6.44	-1
4327	27083	16,17	11	0	11.17	-1
4327	27083	25,26	17	56	-1	3.41
4327	27083	28,29	1	14	-1	11.24
4327	27083	30,31	31	99	-1	2.87
4336	453079	16,17	8	1	8.12	-1
4346	561558	28,29	6	0	7.48	-1
4349	451037	23,24	5	15	-1	2.97
4358	639232	23,24	0	6	-1	5.94
4358	639232	25,26	10	0	9.65	-1
4358	639232	30,31	9	0	10.01	-1
4361	461835	15,16	7	0	7.4	-1
4361	461835	28,29	5	0	6.23	-1
4366	1093	27,28	43	20	2.33	-1
4366	1093	30,31	5	18	-1	3,24
4371	644240	15,17	7	0	7,51	-1
4375	388688	27,29	8	2	5,4	-1
4375	388688	28,29	17	2	10.59	-1
4380	648996	15,16	6	0	6.34	-1
4380	648996	15,17	6	0	6,44	-1
4406	449836	15,17	8	1	8,59	-1
4413	640525	15,16	7	0	7,4	-1
4413	640525	15,17	7	0	7.51	-1
4417	388085	15,16	9	0	9,51	-1
4417	388085	15,17	9	0	9.66	-1
4421	448787	15,17	23	9	2.74	-1
4421	448787	16,17	22	9	2.48	-1
4435	554742	16,17	0	6	-1	5.91
4441	549640	27,28	0	7	-1	6,46
4441	549640	30,31	15	6	2.78	-1
4445	530774	15,17	5	16	-1	2.98
4446	375814	15,16	8	0	8,46	-1
4446	375814	15,17	8	1	8.59	-1
4447	446789	15,16	16	5	3,38	-1
4450	503491	16,17	6	0	6.09	-1
4452	639441	23,24	1	8	-1	7.92
4454	649035	27,28	0	7	-1	6.46
4454	649035	28,29	7	0	8.72	-1
4461	470602	15,16	12	1	12.68	-1
4461	470602	16,17	1	9	-1	8,86
4477	453708	15,17	1	19	-1	17.7
4477	453708	16,17	1	19	-1	18.71
4483	551527	15,17	0	7	-1	6.52
4503	549945	15,16	12	2	6,34	-1
4503	549945	15,17	12	2	6.44	-1
4505	391511	27,28	8	0	8.66	-1

TEO TE VIO	CI TIONED	PAIR	GT 03370G 1	GY 633335	RATIO	RATIO
SEQ ID NO	CLUSTER	AB		CLONES B	PLUS	MINUS
4515	420686	16,17	0	8	-1	7.88
4517	551995	28,29	7	0	8.72	-1
4523	451032	15,16	6	0	6.34	-1
4538	557852	28,29	6	0	7.48	-1
4542	554500	15,16	6	0	6.34	-1
4550	43642	16,17	0	6	-1	5.91
4551	595506	15,16	0	6	-1	5.68
4553	555702	15,16	11	2	5.81	-1
4556	519109	23,24	1	34	-1	33.65
4557	450829	15,17	0	8	-l	7.45
4557	450829	16,17	0	8	-1	7.88
4598	386940	27,28	0	14	-1	12.93
4598	386940	28,29	14	0	17.44	-1
4611	375380	15,17	1	9	-1	8.38
4611	375380	16,17	0	9	-1	8.86
4619	553850	28,29	5	0	6,23	-1
4627	557401	15,17	10	2	5.37	-1
4632	349744	16,17	0	6	-1	5.91
4635	449437	15,16	14	3	4.93	-1
4635	449437	16,17	3	12	-1	3.94
4635	449437	27,28	0	8	-1	7.39
4635	449437	28,29	8	0	9.97	-1
4645	562292	16,17	0	6	-1	5.91
4652	224812	16,17	31	63	-1	2
4652	224812	25,26	200	686	-1	3.55
4652	224812	27,29	84	38	2.98	-1
4652	224812	28,29	80	38	2,62	-1
4670	446739	28,29	5	0	6.23	-1
4684	247	15,17	15	44	-1	2,73
4684	247	18,19	0	15	-1	13.13
4684	247	30,31	11	51	-1	4.17
4686	491799	15,16	0	11	-1	10.41
4691	461	15,16	14	39	-1	2.64
4691	461	15,17	14	34	-1	2.26
4691	461	18,19	1	10	-1	8,75
4691	461	19,20	10	0	7.48	-1
4691	461	27,29	41	124	-1	2,24
4691	461	28,29	37	124	-1	2.69
4691	461	30,31	21	119	-1	5.09
4696	562292	16,17	0	6	-1	5.91
4702	487182	15,17	2	11	-1	5.12
4702	487182	16,17	1	11	-1	10.83
4702	487182	25,26	3	72	-1	24.87
4706	561046	16,17	0	6	-1	5.91
4746	556040	23,24	2	15	-1	7.42
4748	452822	15,16	9	23	-1	2.42
4758	455820	16,17	0	7	-1	6.89
4767	556802	28,29	8	0	9.97	-1

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		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
4771	465446	25,26	26	11	2.28	-1
4826	452775	27,28	1	9	-1	8.31
4827	560868	27,29	6	0	8.09	-1
4835	558534	16,17	0	7	-1	6.89
4850	549591	15,17	2	24	-1	11.18
4850	549591	16,17	7	24	-1	3.38
4885	27586	12,13	7	22	-1	3.04
4885	27586	12,14	7	21	-1	2.86
4885	27586	21,22	9	0	8,85	-1
4885	27586	25,26	17	0	16.41	-1
4885	27586	30,31	24	0	26.69	-1
4888	561558	28,29	6	0	7.48	-1
4900	2435	15,17	0	- 8	-1	7.45
4900	2435	27,28	0	7	-1	6.46
4900	2435	28,29	7	1	8.72	-1
4907	554604	15,17	0	7	-1	6.52
4922	559057	15,17	0	7	-1	6.52
4927	553877	16,17	1	8	-1	7.88
4950	549911	16,17	0	6	-1	5.91
4959	409612	25,26	6	0	5.79	-1
4972	226324	15,17	0	8	-1	7.45
4972	226324	16,17	0	8	-1	7.88
4972	226324	27,29	5	0	6.75	-1
4986	413915	12,13	6	0	6.2	-1
4986	413915	12,14	6	0	6.29	-1
5007	2284	23,24	6	17	-1	2.8
5007	2284	25,26	17	4	4,1	-1
5007	2284	30,31	14	32	-1	2.05
5018	365634	15,16	12	2	6.34	-1
5018	365634	16,17	2	14	-1	6.89
5018	365634	25,26	10	0	9.65	-1
5026	549829	16,17	0	6	-1	5.91
5033	411113	16,17	0	6	-1	5.91
5042	1079	03,04	24	55	-1	2.35
5042	1079	28,29	16	44	-1	2:21
5047	397581	28,29	6	20	-1	2.68
5047	397581	27,29	1	20	-1	14.83
5051	23961	15,16	102	44	2.45	-1
5051	23961	15,17	102	40	2.74	-1
5051	23961	25,26	73	169	-1	2.4
5058	446900	15,17	1	11	-1	10.25
5058	446900	16,17	2	11	-1	5.42
5065	448677	15,16	11	1	11.63	-1
5067	560538	16,17	0	8	-1	7.88
5079	62458	15,16	2	14	-1	6.62
5079	62458	15,17	2	11	-1	5.12
5079	62458	25,26	20	227	-1	11.76
5079	62458	30,31	19	245	-1	11.59

		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
5102	556216	15,17	0	7	-1	6.52
5102	556216	16,17	0	7	-l	6.89
5117	549041	16,17	0	6	-1	5.91
5133	453004	27,28	0	8	-1	7.39
5133	453004	28,29	8	0	9.97	-1
5135	560868	27,29	6	0	8.09	-1
5147	403419	13,14	0	9	-l	8.87
5147	403419	27,29	7	0	9.44	-1
5161	559806	16,17	0	6	-1	5.91
5168	550562	27,28	1	9	-1	8.31
5168	550562	28,29	9	1	11.21	-1
5186	91178	15,17	0	7	-1	6.52
5188	561485	15,17	0	7	-1	6.52
5188	561485	16,17	0	7	-1	6.89
5195	558412	30,31	6	0	6.67	-1
5200	453846	15,17	0	11	-1	10.25
5221	562565	15,16	0	8	-1	7.57
5221	562565	27,28	11	1	11.91	-1
5221	562565	27,29	11	0	14.84	-1
5228	553705	15,16	12	0	12.68	-1
5228	553705	28,29	8	0	9.97	-1
5231	562459	16,17	2	10	-1	4.92
5231	562459	27,28	13	29	-1	2.06
5231	562459	27,29	13	1	17.54	-1
5231	562459	28,29	29	1	36.13	-1
5236	62458	15,16	2	14	-1	6.62
5236	62458	15,17	2	11	-1	5.12
5236	62458	25,26	20	227	-1	11.76
5236	62458	30,31	19	245	-1	11.59
5240	448741	28,29	6	0	7.48	-1
5249	451401	15,17	9	1	9.66	-1
5258	451802	15,17	0	9	-1	8.38
5258	451802	16,17	1	9	-1	8.86
5264	452775	27,28	1	9	-1	8.31
5268	92639	12,14	2	11	-1	5.25
5271	452500	27,28	0	15	-1	13.85
5271	452500	28,29	15	5	3.74	-1
5272	452204	15,16	9	1	9.51	-1
5272	452204	16,17	1	8	-1	7.88
5272	452204	23,24	3	13	-1	4.29
5272	452204	27,28	0	14	-1	12.93
5272	452204	28,29	14	0	17.44	-1
5273	447025	15,16	48	19	2.67	-1
5273	447025	27,28	0	8	-1	7.39
5273	447025	27,29	0	13	-1	9.64
5277	452052	15,17	8	1	8.59	-1
5277	452052	27,28	2	14	-1	6.46
5277	452052	28,29	14	0	17.44	-1

		PAIR	I		RATIO	RATIO
SEO ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
5281	452142	15,16	1	10	-1	9.46
5281	452142	16,17	10	2	5.08	-1
5282	451994	27,28	4	17	-1	3.92
5297	4244	28,29	8	28	-1	2.81
5308	450262	21.22	0	8	-1	8,13
5308	450262	27,28	30	14	2.32	-1
5308	450262	28,29	14	46	-1	2,64
5311	452506	15.16	8	Ti-	8 46	-1
5311	452506	28,29	6	0	7.48	-1
5313	7022	25,26	7	0	6.76	-1
5313	7022	27,28	4	16	-1	3.69
5313	7022	28,29	16	5	3.99	-1
5315	2930	01,02	3	14	-1	5.06
5315	2930	27.29	5	0	6.75	-1
5315	2930	28,29	7	0	8.72	-1
5317	454226	28,29	5	0	6.23	-1
5321	453470	15,16	12	1	12.68	-1
5321	453470	15,17	12	1	12.88	-1
5324	454050	27,28	0	7	-1	6.46
5338	454518	27,29	6	0	8.09	-1
5338	454518	28,29	6	-0	7.48	-1
5350	23649	27,28	37	12	3.34	-1
5350	23649	27,29	37	0	49.92	-1
5350	23649	28,29	12	0	14.95	-1
5357	519109	23,24	1	34	-1	33.65
5360	453783	27,28	0	8	-1	7,39
5360	453783	28,29	8	0	9.97	-1
5365	454509	25,26	7	0	6.76	-1
5365	454509	27,28	6	0	6.5	-1
5366	454562	23,24	13	4	3.28	-1
5366	454562	25,26	7	0	6.76	-1
5370	453783	27,28	0	8	-1	7.39
5370	453783	28,29	8	0	9.97	-1_
5373	801	30,31	25	7	3.97	-1
5374	453494	27,28	7	0	7.58	-1
5374	453494	30,31	6	0	6.67	-1
5375	453202	15,16	6	0	6.34	-1
5375	453202	16,17	0	11	-1	10.83
5381	387530	30,31	2	24	-1	10.79
5382	453846	15,17	0	11	-1	10.25
5391	551995	28,29	7	0	8.72	-1
5397	446531	16,17	0	6	-1	5.91
5401	453508	25,26	27	9	2.9	-1_
5401	453508	27,28	9	1 1	9.75	-1
5413	560868	27,29	6	0	8.09	-1
5448	554742	16,17	0	6	-1	5.91
5467	551617	15,16	9	0	9.51	-1
5467	551617	16,17	0	7	-1	6.89

		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
5489	468672	15,16	6	0	6.34	-1
5489	468672	16,17	0	8	-1	7.88
5501	402836	12,13	130	63	2.13	-1
5501	402836	23,24	17	4	4.29	-1
5508	561707	27,29	6	0	8.09	-1
5508	561707	30,31	9	1	10.01	-1
5509	561180	15,16	11	2	5.81	-1
5513	562137	23,24	8	0	8,08	-1
5518	449356	15,17	7	21	-1	2.79
5518	449356	16,17	4	21	-1	5.17
5525	549511	15,17	0	7	-1	6.52
5525	549511	16,17	0	7	-1	6.89
5526	560700	16,17	0	6	-1	5.91
5535	452523	15,17	10	2	5.37	-1
5537	406092	18,19	16	2	9.14	-1
5537	406092	25,26	7	0	6.76	-1
5537	406092	27,28	1	26	-1	24.01
5537	406092	27,29	1	28	-1	20.76
5537	406092	30,31	13	4	3.61	-1
5542	235874	12,13	62	239	-1	3.73
5542	235874	12,14	62	317	-1	4.88
5542	235874	25,26	102	21	4.69	-1
5542	235874	27,28	73	16	4.94	-1
5542	235874	28,29	16	70	-1	3.51
5542	235874	30,31	119	9	14.71	-1
5550	452957	27,29	5	0	6.75	-1
5550	452957	28,29	6	0	7.48	-1
5554	455855	27,29	8	0	10.79	-1
5559	510254	15,16	17	1	17.97	-1
5559	510254	16,17	1	8	-1	7.88
5560	451812	27,29	7	1	9.44	-1
5565	450225	15,17	11	3	3.94	-1
5565	450225	27,28	1	24	-1	22.16
5565	450225	28,29	24	1	29.9	-1
5572	503122	25,26	8	178	-1	23.05
5572	503122	27,29	12	2	8.09	-1
5572	503122	30,31	22	98	-1	4
5578	446936	23,24	10	26	-1	2.57
5578	446936	25,26	17	38	-1	2.32
5578	446936	27,28	158	31	5.52	-1
5578	446936	27,29	158	27	7.89	-1
5589	453079	16,17	8	1	8.12	-1
5599	454226	28,29	5	0	6.23	-1
5600	452973	27,28	0	9	-1	8.31
5600	452973	28,29	9	2	5.61	-1
5606	452648	23,24	2	15	-1	7.42
5611	366607	27,28	1	12	-1	11.08
5611	366607	28,29	12	0 -	14.95	-1

		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
5617	521840	15,16	6	0	6.34	-1
5620	523674	28,29	5	0	6.23	-1
5624	452775	27,28	1	9	-1	8.31
5634	454460	27,28	0	13	-1	12.01
5634	454460	28,29	13	2	8.1	-1
5662	456001	28,29	10	1	12.46	-1
5665	450914	15,16	0	6	-1	5.68
5666	455855	27,29	8	0	10.79	-1
5688	400258	16,17	6	0	6.09	-1
5688	400258	27,28	7	.0	7.58	-1
5697	2544	01,02	23	9	2.36	-l
5697	2544	03,04	13	30	-1	2.37
5699	402534	12,13	8	21	-1	2.54
5704	456567	28,29	5	0	6.23	-1
5710	455601	15,17	10	0	10.73	-1
5710	455601	16,17	8	0	8.12	-1
5712	402916	23,24	8	0	8.08	-1
5712	402916	27,28	22	5	4.76	-1
5712	402916	27.29	22	4	7.42	-1
5714	27586	12.13	7	22	-1	3.04
5714	27586	12,14	7	21	-1	2.86
5714	27586	21,22	9	0	8.85	-1
5714	27586	25.26	17	0	16.41	-1
5714	27586	30,31	24	0	26.69	-1
5732	407711	15,16	0	6	-1	5.68
5733	411113	16.17	0	6	-1	5.91
5742	400426	12,14	22	10	2.31	-1
5752	155374	15.16	6	0	6.34	-1
5752	155374	15,17	6	0	6.44	-1
5752	155374	27,28	27	i	29.24	-1
5752	155374	27,29	27	0	36.42	-1
5753	409612	25.26	6	0	5.79	-1
5759	1054	03,04	49	23	2.08	-1
5759	1054	12,14	54	24	2.36	-1
5759	1054	15.17	64	28	2.45	-1
5759	1054	18,20	3	19	-1	7.41
5759	1054	19,20	5	19	-1	5.08
5759	1054	27,29	64	31	2.79	-1
5759	1054	28.29	69	31	2.77	-1
5759	1054	30,31	27	15	2	-1
5761	400454	12,14	9	1	9.43	-1
5765	409589	27,29	5	0	6.75	-1
5768	20517	12,13	18	3	6.2	-1
5768	20517	13,14	3	18	-1	5,91
5769	413915	12,13	6	0	6.2	-1
5769	413915	12,14	6	0	6.29	-1
5772	401510	12,14	8	0	8.39	-1
5775	417259	15,17	4	14	-1	3,26

		PAIR			RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
5775	417259	16,17	1	14	-1	13.79
5776	20517	12,13	18	3	6.2	-1
5776	20517	13,14	3	18	-1	5.91
5778	402534	12,13	8	21	-1	2.54
5780	419255	15,16	11	0	11,63	-1
5780	419255	15,17	11	1	11.81	-1
5788	402070	27,28	1	9	-1	8.31
5788	402070	28,29	9	0	11.21	-1
5789	402070	27,28	1	9	-1	8.31
5789	402070	28,29	9	0	11.21	-1
5792	402353	12,13	13	4	3.36	-1
5794	163970	18,20	12	25	-1	2.44
5794	163970	19,20	7	25	-1	4.77
5794	163970	27,28	7	39	-1	5.15
5794	163970	28,29	39	7	6.94	-1
5794	163970	30,31	29	6	5.38	-1
5801	268336	15,16	8	1	8.46	-1
5807	20517	12,13	18	3	6.2	-1
5807	20517	13,14	3	18	-1	5.91
5822	420686	16,17	0	8	-1	7.88
5823	447579	23,24	3	13	-1	4.29
5823	447579	28,29	7	0	8.72	-1
5825	463824	15,16	6	0	6.34	-1
5825	463824	15,17	6	0	6.44	-1
5825	463824	27,28	0	8	-1	7.39
5825	463824	28,29	8	0	9.97	-1
5827	388688	27,29	- 8	2	5.4	-1
5827	388688	28,29	17	2	10.59	-1
5840	446964	15,17	1	12	-1	11.18
5860	403949	12,13	11	1	11.36	-1
5862	235874	12,13	62	239	-1	3.73
5862	235874	12,14	62	317	-1	4.88
5862	235874	25,26	102	21	4.69	-1
5862	235874	27,28	73	16	4.94	-1
5862	235874	28,29	16	70 .	-1	3.51
5862	235874	30,31	119	9	14.71	-1
5876	644342	27,29	6	0	8.09	-1
5897	644342	27,29	6	0	8.09	-1
5898	419479	23,24	2	11	-1	5.44
5906	451508	30,31	7	0	7.79	-1
5924	218416	16,17	0	6	-1	5.91
5933	463824	15,16	6	0	6.34	-1
5933	463824	15,17	6	0	6.44	-1
5933	463824	27,28	0	8	-1	7.39
5933	463824	28,29	8	0	9.97	-1
5934	389377	27,28	7	0	7.58	-1
5934	389377	30,31	10	1	11.12	-1
5943	18786	15,17	6	0	6,44	-1

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		PAIR		1 1	RATIO	RATIO
SEQ ID NO	CLUSTER	AB	CLONES A	CLONES B	PLUS	MINUS
5943	18786	30,31	6	0	6.67	-l
5944	447494	15,16	26	8	3.44	-1
5966	1454	03,04	38	14	2.65	-1
5966	1454	12,13	37	15	2.55	-1
5966	1454	13,14	15	35	-1	2,3
5966	1454	25,26	12	3	3.86	-1
5966	1454	30,31	7	22	-1	2.83
5969	15296	27,28	6	0	6.5	-1
5979	446673	16,17	8	1	8.12	-1
5991	446341	28,29	6	0	7.48	-1
6002	379335	15,16	11	0	11.63	-1
6002	379335	15,17	11	0	11.81	-1
6003	228873	27,29	20	0	26.98	-1
6003	228873	28,29	14	0	17.44	-1
6004	446663	15,17	14	32	-1	2.13
6004	446663	16,17	6	32	-1	5.25
6004	446663	18,20	1	8	-1	9,36
6004	446663	19,20	0	8	-1	10.7

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Comment	invasive adenocarcinom a, moderately differentiated; focal perineural invasion is seen	Hyperplastic polyp in appendix.	Perineural invasion; donut anastomosis Neg. One tubulovillous and one tubular adenoma with no high grade dvsplasia.	patient history of metastatic melanoma	
Dist Met Grade	MX	MO	MO	MO	WO
Dist Met & Loc	Neg	Neg	Neg	Neg	Neg
Reg Lymph Grade	Z	NO N	0 N	0N	Z
Lymph Met Incid	3/8	0/12	0/34	61/0	1/5
Lymph Lymph Met Met Incid	Pos	Neg	Neg	Neg	Pos
Local Invasion	Extending into subserosal adipose tissue	Invasion through muscularis propria, subserosal involvement; ileocec. valve involvement	Invasion of muscularis propria into serosa, into serosa, survolving submucosa of urinary bladder	Invasion through the muscularis propria into suscrosal adipose tissue. Heocecal junction.	Invasion of muscularis propria into percolonic fat
Histo Grade	<b>G</b> 2	8	G5	E5	G
Size Grade Histo	E	E	T4	13	T3
Size	4.0	9.0	9	9	5.0
Anatom	Ascending	Cecum	Sigmoid	Cecum	Transverse colon
Стр	Ш	Ħ	Ħ	=	Ħ
Path	21	71	140	44	147
PtD	15	52	121	125	128

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Comment					Small separate	tubular	adenoma (0.4	cm)					Perineural	invasion	identified	adjacent to	metastatic	adenocarcinom	ę	Separate	tubolovillous	and tubular	adenomas					
Dist	Met Grade	M			Mo Mo						,		M							M0								
	& Loc	Neg			Neg								Pos-	Liver						Neg								
Reg	Lymph Grade	NZ			0 N								N2							Z								
$\vdash$		10/24			6/0								7/21							2/13								
Lymph	Mct	Pos			Neg								Pos							Pos								
Local Invasion		through wall and	into surrounding	adipose tissue	Invasion through	muscularis propria	into non-	peritonealized	pericolic tissue;	gross	configuration is	annular.	Invasion of	muscularis propria	into pericolonic	adipose tissue, but	not through serosa.	Arising from	tubular adenoma.	Invasion through	mucsularis propria	into	subserosa/pericolic	adipose, no serosal	involvement.	Gross	configuration	annular.
Histo	Grade				3								G5							G5								
Size Grade Histo		E			13								T3							13								
Size		5.5			5.0					_			5.5							3.8								
a	Loc	Splenic	flexure		Rectum								Cecum							Hepatic	flexure							
ф					=								Z							Ħ								
Path	8	149			152								160				_			175								
Pt ID		130			133								141							156								_

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Comment	Hyperplastic polyps	Tubulovillous adenoma with high grade dysplasia			Descending colon polyps, no HGD or carcinoma identified
Dist Met Grade	MX	M0	MX	MO	Mo
Dist Met & Loc	Neg	Neg	Pos - Mesente ric deposit	Neg	Neg
Reg Lymph Grade	IN	0N	IN	N0	N2
Lymph Met Incid	1/8	01/0	0/15	0/12	7/10
Lymph Met	Pos	Neg	Neg	Neg	Pos
Local Invasion	G2 to Invasion through G3 muscularis propria to involve subserosal, perirectoal adipose, and serosa	Invasion through muscularis propria into subserosal adipose tissue.	Invades through muscularis propria to involve pericolonic adipose, extends to serosa.	Invades full thickness of muscularis propria, but mesenteric adipose free of malignancy	Invasion into perirectal adipose tissue.
Size Grade Histo	G2 to	G2	G2	G2	G2
Grade	EI	T3	T3	72	T3
Size	5.8	5.5	6	6.5	4
Anatom	Rectum	Ascending	Transverse	Cecum	Rectum
ģ	Ħ	=	Ħ	-	Ħ
Path D	247	283	285	287	297
Pt ID	228	264	266	268	278

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Comment	Tubulovillous adenoma (2.0 cm) with no high grade dysplasia. Neg. liver biopsy.	1 hyperplastic polyp identified			Two mucosal polyps	Tumor arising at prior ileocolic surgical anastomosis.
Dist Met Grade	W0	. W0	MX	MO	MO	MI
Dist Met & Loc	Neg	Neg	Neg	Neg	Neg	Pos - Liver
Reg Lymph Grade	z	0X	0X	0 Z	Z	Z
Lymph Met Incid	2/12	9/0	0/4	0/4	1/5	1/6
Lymph	Pos	See.	Neg	Neg	Pos	Pos
Local Invasion	Invasion through muscularis propria and invades pericolic adipose tissue. Heocecal junction.	Extends into perirectal fat but does not reach scrosa	Invasion through muscularis propria to involve pericolonic fat. Arising from villous adenoma.	Through colon wall into subserosal adipose tissue. No serosal spread seen.	Invasion thru muscularis propria to pericolonic fat	Invasion through muscularis propria into subserosal adipose tissue, not serosa.
Histo Grade	B	25	B	25	25	62
Size Grade Histo	2	E E	E	T	E	T3
Size	5.5	9	2 cm inva sivc	6.5	4.3	2
Anatom	Cecum	Rectosigm	Ascending	Sigmoid	Ascending	Ascending
dy.	Ħ	Ħ	Ħ	Ħ	Ħ	2
Path	315	358	360	375	412	444
PtID	296	339	341	356	360	392

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Comment		rediagnosis of oophorectomy path to metastatic colon cancer.	Anatomical location of primary not notated in report. Evidence of chronic colitis.	No mention of distant met in report
Dist Met Grade	MO	M1	M	M0
Dist Met Dist & Loc Met Grade	Neg	Pos - Liver	Pos - Liver	SeN
Reg Lymph Grade	0X	0 N	ĭ	N2
Lymph Met Incid	0/21	2/0	2/17	9/9
Lymph Met	Neg	Neg	Pos	Pos
Local Invasion Lymph Lymph Met Met Met Incid	Cecum, invades through muscularis propria to involve subserosal adipose tissue but not serosa.	Invasive through muscularis to involve periserosal fat; abutting ileocecal junction.	Invasion through muscularis propria involving pericolic adipose, serosal surface uninvolved	penetrates muscularis propria, involves pericolonic fat.
Size Grade Histo	8	S	8	G2
Grade	E	E	E	E
Size	0.9	8.	7.5	m
Anatom Loc	Cecum	Cecum		Sigmoid
Сф	П	Z	N	≥i
Path	445	465	383	395
Pt ID	393	413	505	517

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Table 6

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Comment	Omentum with fibrosis and fat necrosis. Small bowel with acute and chronic serrositis, focal abscess and adhesions.		Appendix dilated and fibrotic, but not involved by tumor
Dist Met Grade	MO	M	MO
Dist Met Dist & Loc Met Grade	Neg	Pos - Liver	Neg
Reg Lymph Grade	NO N	N2	NO NO
Lymph Met Incid	8/0	6/12	85/0
Lymph Lymph Met Met Incid	Neg	Pos	Neg
Local Invasion	Invasion through the muscularis propriate involving pericolic fat. Serosa free of tumor.	Invasion through muscularis propria extensively through submucosal and extending to serosa.	Invasion through the bowel wall, into suberosal adipose. Serosal surface free of tumor.
Histo Grade	8		G2
Size Grade Histo	T3	T3	T3
	21	5.5	11.5
Anatom	Ascending colon	Ascending 5.5 colon	Cecum
Ę.	Ħ	ΣI	П
Path ID	553	565	596
Pt ID	534	546	577

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Comment	moderately differentiated adenocarcinom a with mucinous differentiation (% not stated), tubular adenoma and hyperplstic polyps present,	invasive poorly differentiated adenosquamous carcinoma	moderately differentiated invasive adenocarcinom a	Peritumoral lymphocytic response; 5 LN examined in pericolic fat, no metastatases observed.	Three fungating lesions examined.
Dist Met Grade	MX	M	M	MX	M
Dist Met & Loc	Neg	Pos - Liver	Pos - Liver	Neg	Pos - Liver
Reg Lymph Grade	N N	N2	N0	0N	N2
Lymph Met Incid	0/22	5/17	0/12		5/10
Lymph Met	Neg	Pos	Neg	Neg 8	Pos
Local Invasion	extending through bowel wall into serosal fat	through muscularis propria into pericolic soft tissues	through muscularis propria into pericolic fat, but not at serosal surface	G2-G3 Invasion of muscularis propria into soft tissue	G2-G3 Extending through muscularis propria into pericolonic fat
Histo Grade	G2	8	92	G2-G3	G2-G3
Size Grade	T3	E	T3	E	T3
Size	14.0	3.5	5.6	2.5	5.0
Anatom	Cecum	Ascending	Descendin g colon	Rectosigm	Cecum
Grp	Ħ	≥i	Δ	н	IV
Path ID	714	803	508	908	808
Pt ID	695	784	786	787	789

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Comment		poorly	differentiated invasive colonic	adenocarcinom	well to moderately	differentiated	adenocarcinom	as; this patient	has tumors of	the ascending	colon and the	sigmoid colon	moderately	differentiated	adenocarcinom	ra Tables			Perineural	invasion	present.
Dist Met Grade	M	M			MI								M				M		MI		
Dist Met & Loc	Pos - Liver	Pos-	Liver		Pos - Liver								Pos-	Liver		,	Fos-	Liver	Pos -	Liver	
Reg Lymph Grade	Į.	N2			N0								Z				N7		N2		
Lymph Met Incid	3/13	13/25			3/21								74			2.1.	CI/II		4/15		
Lymph Lymph Met Met Incid	Pos	Pos			Pos								Pos			,	S		Pos		
Local Invasion	G1-G2 Invading through muscularis propria into perirectal fat	Through the	muscularis propria into pericolic fat		Into muscularis propria	•							Through	muscularis propria	int subscrosal	tissue	I brough	muscularis propria	Invasion through	muscularis propria	into perirectal soft tissue
Histo Grade	G1-G2	g			C1								3			Т	3		GZ		
Grade	E	T3			TZ								13			i	5		T3		
Size	8.9	5.00			2.0								8.			1			5.2		
Anatom Size Grade Histo Loc Grade Grade	Rectum	Ascending	colon		Ascending								Cecum				Ascending	colon	Rectum		
Сър	N	2			Ŋ	-			_				2				≥		N		
Path	608	810			806								606			1	910		911		
Pt ID	790	161			888								886			-	268		891		

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Comment	Perineural invasion present, extensive. Patient with a history of colon cancer.	Perineural invasion focally present. Omentum mass, but resection with no tumor identified.	M1 Primary adenocarcinom a arising from tubulovillous adenoma.
Dist Met Grade	MI	MI	MI
Dist Met & Loc	Pos - Liver, left and right lobe, omentu m	Pos - Liver	Pos - Liver
Reg Lymph Grade	ĸ	NZ	ī
Lymph Met Incid	1/28	14/17	1/1
Lymph Met	Pos	Pos	Pos
Local Invasion Lymph Lymph Mct Mct Incid	Invasion into pericolic sort tissue. Tumor focally invading skeletal muscle attached to colon.	G2-G3 Through muscularis propria into pericolic fat	G2 Invasion through colon wall and focally involving subserosal tissue.
Histo Grade	62	G2-G3	G2
Grade	EL	EL	T3
Size	5.0	0.9	0.9
Anatom Size Grade Histo Loc . Grade	Sigmoid	colon	Sigmoid
Стр	2	2	IV
Path	912	913	1009
Pt D Path	892	893	686

Table 8

SEQ ID NO         SpotID         T/N Colon >2x         T/N Colon <16x	Tabl	e 8			
6 43971 0.0 75.0 8.0 31 40453 0.0 42.9 7.0 40 40457 0.0 71.4 7.0 44 46308 0.0 50.0 8.0 50 45610 0.0 62.5 8.0 50 45610 0.0 50.0 8.0 72 44673 0.0 50.0 8.0 72 44673 0.0 50.0 8.0 74 42422 0.0 37.5 8.0 81 44679 0.0 50.0 8.0 81 44679 0.0 50.0 8.0 81 44679 0.0 50.0 8.0 81 44679 0.0 50.0 8.0 81 44679 0.0 50.0 8.0 81 44679 0.0 50.0 8.0 81 44679 0.0 50.0 8.0 81 44679 0.0 50.0 8.0 81 44679 0.0 50.0 8.0 81 44218 0.0 37.5 8.0 81 44679 0.0 50.0 8.0 81 44216 0.0 57.5 8.0 133 39755 0.0 42.9 7.0 152 44216 0.0 37.5 8.0 153 38367 0.0 42.9 7.0 166 43508 0.0 37.5 8.0 161 41869 0.0 42.9 7.0 161 41869 0.0 42.9 7.0 162 43508 0.0 37.5 8.0 164 338357 0.0 57.1 7.0 166 39069 0.0 42.9 7.0 167 39061 0.0 57.1 7.0 180 39767 0.0 42.9 7.0 184 43881 0.0 37.5 8.0 186 43873 0.0 37.5 8.0 186 43873 0.0 37.5 8.0 197 46330 0.0 37.5 8.0 198 43471 0.0 57.1 7.0 197 46330 0.0 37.5 8.0 198 43471 0.0 57.1 7.0 197 46330 0.0 37.5 8.0 198 42479 0.0 50.0 8.0 202 42479 0.0 50.0 8.0 218 46015 0.0 42.9 7.0 217 46007 0.0 42.9 7.0 228 45301 0.0 37.5 8.0 228 45301 0.0 37.5 8.0 228 45301 0.0 37.5 8.0 228 45301 0.0 37.5 8.0 228 45301 0.0 37.5 8.0 228 45301 0.0 37.5 8.0 229 7.0 24279 7.0 50.0 50.0 8.0 2429 7.0 24279 7.0 50.0 57.1 7.0 2568 41027 0.0 42.9 7.0 274 41737 0.0 42.9 7.0 274 41737 0.0 42.9 7.0 274 41737 0.0 42.9 7.0 274 41737 0.0 42.9 7.0	SEO ID NO	SnotID	T/N Colon >2v	T/N Colon Aple	T/N Colon Name Bation
31         40453         0.0         42.9         7.0           40         40457         0.0         71.4         7.0           44         446308         0.0         50.0         8.0           50         45610         0.0         62.5         8.0           70         42816         0.0         50.0         8.0           72         44673         0.0         50.0         8.0           74         42422         0.0         37.5         8.0           81         44679         0.0         50.0         8.0           84         42418         0.0         37.5         8.0           83         133         39755         0.0         42.9         7.0           139         44916         0.0         50.0         8.0           147         45618         0.0         37.5         8.0           149         44926         0.0         50.0         8.0           153         38367         0.0         42.9         7.0           153         38357         0.0         57.1         7.0           161         41869         0.0         37.5         8.0					
40					
44         46308         0.0         50.0         8.0           50         45610         0.0         62.5         8.0           70         42816         0.0         50.0         8.0           72         44673         0.0         50.0         8.0           74         42422         0.0         37.5         8.0           81         44679         0.0         50.0         8.0           84         42418         0.0         37.5         8.0           133         39755         0.0         42.9         7.0           139         44916         0.0         50.0         8.0           147         45618         0.0         37.5         8.0           152         44216         0.0         37.5         8.0           153         38367         0.0         42.9         7.0           153         38367         0.0         42.9         7.0           153         38367         0.0         42.9         7.0           161         41869         0.0         37.5         8.0           162         43508         0.0         37.5         8.0 <t< td=""><td></td><td></td><td></td><td></td><td></td></t<>					
50         45610         0.0         62.5         8.0           70         42816         0.0         50.0         8.0           72         44673         0.0         50.0         8.0           74         42422         0.0         37.5         8.0           77         43983         0.0         37.5         8.0           81         44679         0.0         50.0         8.0           84         42418         0.0         37.5         8.0           133         39755         0.0         42.9         7.0           139         44916         0.0         50.0         8.0           147         45618         0.0         37.5         8.0           152         44216         0.0         37.5         8.0           153         38367         0.0         42.9         7.0           153         38367         0.0         57.1         7.0           161         41869         0.0         37.5         8.0           162         43508         0.0         37.5         8.0           164         38365         0.0         57.1         7.0 <t< td=""><td></td><td></td><td></td><td></td><td></td></t<>					
70         42816         0.0         50.0         8.0           72         44673         0.0         50.0         8.0           74         442422         0.0         37.5         8.0           77         43983         0.0         37.5         8.0           81         44679         0.0         50.0         8.0           34         42418         0.0         37.5         8.0           133         39755         0.0         42.9         7.0           139         44916         0.0         50.0         8.0           147         45618         0.0         37.5         8.0           152         44216         0.0         37.5         8.0           152         44216         0.0         37.5         8.0           153         33357         0.0         57.1         7.0           161         41869         0.0         42.9         7.0           161         44850         0.0         37.5         8.0           164         38365         0.0         57.1         7.0           166         39069         0.0         42.9         7.0					
72         44673         0.0         50.0         8.0           74         42422         0.0         37.5         8.0           77         43983         0.0         37.5         8.0           81         44679         0.0         50.0         8.0           84         42418         0.0         37.5         8.0           133         39755         0.0         42.9         7.0           139         44916         0.0         50.0         8.0           147         45618         0.0         37.5         8.0           149         44926         0.0         50.0         8.0           152         44216         0.0         37.5         8.0           153         38367         0.0         42.9         7.0           161         41869         0.0         42.9         7.0           161         41869         0.0         37.5         8.0           164         43508         0.0         37.5         8.0           166         39069         0.0         42.9         7.0           171         39061         0.0         57.1         7.0					
74         42422         0.0         37.5         8.0           77         43983         0.0         37.5         8.0           81         44679         0.0         50.0         8.0           84         42418         0.0         37.5         8.0           133         39755         0.0         42.9         7.0           139         44916         0.0         50.0         8.0           147         45618         0.0         37.5         8.0           149         44926         0.0         50.0         8.0           152         44216         0.0         37.5         8.0           153         38367         0.0         42.9         7.0           153         38357         0.0         57.1         7.0           161         41869         0.0         37.5         8.0           164         38365         0.0         57.1         7.0           164         38365         0.0         57.1         7.0           171         39061         0.0         57.1         7.0           180         39767         0.0         42.9         7.0					
77         43983         0.0         37.5         8.0           81         44679         0.0         50.0         8.0           84         442418         0.0         37.5         8.0           133         39755         0.0         42.9         7.0           139         44916         0.0         50.0         8.0           147         45618         0.0         37.5         8.0           152         44216         0.0         37.5         8.0           152         44216         0.0         37.5         8.0           153         38367         0.0         42.9         7.0           161         41869         0.0         42.9         7.0           161         41869         0.0         42.9         7.0           164         38365         0.0         57.1         7.0           166         39069         0.0         42.9         7.0           171         39061         0.0         57.1         7.0           184         43881         0.0         37.5         8.0           184         43881         0.0         37.5         8.0					
81         44(79)         0.0         50.0         8.0           84         42418         0.0         37.5         8.0           133         39755         0.0         42.9         7.0           139         44916         0.0         50.0         8.0           147         45618         0.0         37.5         8.0           149         44926         0.0         50.0         8.0           152         44216         0.0         37.5         8.0           153         38367         0.0         42.9         7.0           151         383857         0.0         42.9         7.0           161         41869         0.0         42.9         7.0           162         43508         0.0         37.5         8.0           164         38365         0.0         37.5         8.0           166         39069         0.0         42.9         7.0           171         39061         0.0         57.1         7.0           184         43811         0.0         37.5         8.0           185         39769         0.0         37.5         8.0					
84         42418         0.0         37.5         8.0           133         39755         0.0         42.9         7.0           139         44916         0.0         50.0         8.0           147         45618         0.0         37.5         8.0           149         44926         0.0         50.0         8.0           152         44216         0.0         37.5         8.0           153         38367         0.0         42.9         7.0           158         38357         0.0         57.1         7.0           161         41869         0.0         42.9         7.0           162         43508         0.0         37.5         8.0           164         38365         0.0         57.1         7.0           164         38365         0.0         57.1         7.0           171         39061         0.0         57.1         7.0           180         39767         0.0         42.9         7.0           184         43813         0.0         37.5         8.0           195         39759         0.0         37.5         8.0					
133         39755         0.0         42.9         7.0           139         44916         0.0         50.0         8.0           147         45618         0.0         37.5         8.0           149         44926         0.0         50.0         8.0           152         44216         0.0         37.5         8.0           153         33357         0.0         42.9         7.0           153         33357         0.0         57.1         7.0           161         44869         0.0         42.9         7.0           162         43508         0.0         37.5         8.0           164         33365         0.0         57.1         7.0           166         39069         0.0         42.9         7.0           171         39061         0.0         57.1         7.0           180         39767         0.0         42.9         7.0           184         43831         0.0         37.5         8.0           195         39769         0.0         57.1         7.0           196         39775         0.0         57.1         7.0					
139         44916         0.0         50.0         8.0           147         45618         0.0         37.5         8.0           149         44926         0.0         50.0         8.0           152         44216         0.0         37.5         8.0           153         38367         0.0         42.9         7.0           151         383657         0.0         57.1         7.0           161         41869         0.0         42.9         7.0           162         43508         0.0         37.5         8.0           164         38365         0.0         57.1         7.0           166         39069         0.0         42.9         7.0           171         39061         0.0         57.1         7.0           180         39767         0.0         42.9         7.0           184         43881         0.0         37.5         8.0           195         39759         0.0         37.5         8.0           195         39775         0.0         57.1         7.0           196         39775         0.0         57.1         7.0					
147         45618         0.0         37.5         8.0           149         44926         0.0         50.0         8.0           152         44216         0.0         37.5         8.0           153         38367         0.0         42.9         7.0           158         38357         0.0         57.1         7.0           161         41869         0.0         42.9         7.0           162         43508         0.0         37.5         8.0           164         38365         0.0         57.1         7.0           166         39069         0.0         42.9         7.0           171         39061         0.0         57.1         7.0           180         39767         0.0         42.9         7.0           184         438381         0.0         37.5         8.0           195         39769         0.0         57.1         7.0           196         39775         0.0         57.1         7.0           197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0					
149         44926         0.0         50.0         8.0           152         44216         0.0         37.5         8.0           153         38367         0.0         42.9         7.0           158         38357         0.0         57.1         7.0           161         41869         0.0         42.9         7.0           162         43508         0.0         37.5         8.0           164         38365         0.0         57.1         7.0           166         39069         0.0         42.9         7.0           171         39061         0.0         57.1         7.0           180         39767         0.0         42.9         7.0           184         43831         0.0         37.5         8.0           195         39769         0.0         57.1         7.0           196         39775         0.0         57.1         7.0           196         39775         0.0         57.1         7.0           197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0					
152         44216         0.0         37.5         8.0           153         38367         0.0         42.9         7.0           158         38367         0.0         42.9         7.0           161         41869         0.0         42.9         7.0           162         43508         0.0         37.5         8.0           164         38365         0.0         57.1         7.0           166         39069         0.0         42.9         7.0           171         39061         0.0         57.1         7.0           180         39767         0.0         42.9         7.0           184         43881         0.0         37.5         8.0           195         39759         0.0         57.1         7.0           196         39775         0.0         57.1         7.0           197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0           200         41173         0.0         42.9         7.0           217         46007         0.0         50.0         8.0					
153         38367         0.0         42.9         7.0           158         38357         0.0         57.1         7.0           161         41869         0.0         42.9         7.0           162         43508         0.0         37.5         8.0           164         38365         0.0         57.1         7.0           166         39069         0.0         42.9         7.0           171         39061         0.0         57.1         7.0           180         39767         0.0         42.9         7.0           184         43881         0.0         37.5         8.0           195         39769         0.0         57.1         7.0           196         39775         0.0         57.1         7.0           197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0           200         41173         0.0         37.5         8.0           201         42479         0.0         50.0         8.0           202         42479         0.0         50.0         8.0					
158         38357         0.0         57.1         7.0           161         41869         0.0         42.9         7.0           162         43508         0.0         37.5         8.0           164         38365         0.0         57.1         7.0           166         39069         0.0         42.9         7.0           171         39061         0.0         57.1         7.0           180         39767         0.0         42.9         7.0           184         43831         0.0         37.5         8.0           195         39769         0.0         57.1         7.0           196         39775         0.0         57.1         7.0           197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0           200         41173         0.0         42.9         7.0           202         42479         0.0         50.0         8.0           216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0					
161         41869         0.0         42.9         7.0           162         43508         0.0         37.5         8.0           164         38365         0.0         57.1         7.0           166         39069         0.0         42.9         7.0           171         39061         0.0         57.1         7.0           180         39767         0.0         42.9         7.0           184         43881         0.0         37.5         8.0           186         43873         0.0         37.5         8.0           195         39775         0.0         57.1         7.0           196         39775         0.0         57.1         7.0           197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0           200         41173         0.0         42.9         7.0           202         42479         0.0         50.0         8.0           216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0					
162         43508         0.0         37.5         8.0           164         38365         0.0         57.1         7.0           166         39069         0.0         42.9         7.0           171         39061         0.0         57.1         7.0           180         39767         0.0         42.9         7.0           184         43881         0.0         37.5         8.0           186         43873         0.0         37.5         8.0           195         39769         0.0         57.1         7.0           196         39775         0.0         57.1         7.0           197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0           200         41173         0.0         42.9         7.0           201         42479         0.0         50.0         8.0           216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0           225         45301         0.0         37.5         8.0					
164         38365         0.0         57.1         7.0           166         39069         0.0         42.9         7.0           171         39061         0.0         57.1         7.0           180         39767         0.0         42.9         7.0           184         43881         0.0         37.5         8.0           185         39769         0.0         57.1         7.0           196         39775         0.0         57.1         7.0           197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0           200         41173         0.0         42.9         7.0           202         42479         0.0         50.0         8.0           216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0           225         45301         0.0         37.5         8.0           225         45301         0.0         37.5         8.0           225         45303         0.0         37.5         8.0					
166         39069         0.0         42.9         7.0           171         39061         0.0         57.1         7.0           180         39767         0.0         42.9         7.0           184         43881         0.0         37.5         8.0           186         43873         0.0         37.5         8.0           195         39775         0.0         57.1         7.0           197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0           200         41173         0.0         42.9         7.0           202         42479         0.0         50.0         8.0           216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0           218         46015         0.0         62.5         8.0           228         45301         0.0         37.5         8.0           228         45303         0.0         37.5         8.0           225         43301         0.0         57.5         8.0					
171         39061         0.0         57.1         7.0           180         39767         0.0         42.9         7.0           184         43881         0.0         37.5         8.0           186         43873         0.0         37.5         8.0           195         39769         0.0         57.1         7.0           196         39775         0.0         57.1         7.0           197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0           200         41173         0.0         42.9         7.0           202         42479         0.0         50.0         8.0           216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0           218         46015         0.0         62.5         8.0           225         45301         0.0         37.5         8.0           228         45303         0.0         37.5         8.0           228         45303         0.0         37.5         8.0					
180         39767         0.0         42.9         7.0           184         43881         0.0         37.5         8.0           186         43873         0.0         37.5         8.0           195         39769         0.0         57.1         7.0           196         39775         0.0         57.1         7.0           197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0           200         41173         0.0         42.9         7.0           202         242479         0.0         50.0         8.0           216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0           218         46015         0.0         62.5         8.0           225         45301         0.0         37.5         8.0           228         45303         0.0         37.5         8.0           225         45301         0.0         57.1         7.0           260         41033         0.0         57.1         7.0					
184         43881         0.0         37.5         8.0           186         43873         0.0         37.5         8.0           195         39769         0.0         57.1         7.0           196         39775         0.0         57.1         7.0           197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0           200         41173         0.0         42.9         7.0           202         42479         0.0         50.0         8.0           216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0           218         46015         0.0         62.5         8.0           225         45301         0.0         37.5         8.0           228         45303         0.0         37.5         8.0           250         41033         0.0         57.1         7.0           260         41027         0.0         42.9         7.0           268         41027         0.0         42.9         7.0					
186         43873         0.0         37.5         8.0           195         39769         0.0         57.1         7.0           196         39775         0.0         57.1         7.0           197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0           200         41173         0.0         42.9         7.0           202         242479         0.0         50.0         8.0           216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0           218         46015         0.0         62.5         8.0           225         45301         0.0         37.5         8.0           228         45303         0.0         37.5         8.0           250         41033         0.0         57.1         7.0           260         41035         0.0         57.1         7.0           268         41027         0.0         42.9         7.0           274         41737         0.0         42.9         7.0					
195         39769         0.0         57.1         7.0           196         39775         0.0         57.1         7.0           197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0           200         41173         0.0         42.9         7.0           202         242479         0.0         50.0         8.0           216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0           218         46015         0.0         62.5         8.0           225         45301         0.0         37.5         8.0           228         45303         0.0         37.5         8.0           250         41033         0.0         57.1         7.0           260         41025         0.0         57.1         7.0           268         41027         0.0         42.9         7.0           274         41737         0.0         42.9         7.0           301         39647         0.0         42.9         7.0					
196   39775   0.0   57.1   7.0     197   46330   0.0   37.5   8.0     198   42471   0.0   37.5   8.0     200   41173   0.0   42.9   7.0     202   42479   0.0   50.0   8.0     216   39621   0.0   42.9   7.0     217   46007   0.0   50.0   8.0     218   46015   0.0   62.5   8.0     225   45301   0.0   37.5   8.0     228   45303   0.0   37.5   8.0     228   45303   0.0   37.5   8.0     229   45304   0.0   57.1   7.0     260   41035   0.0   57.1   7.0     268   41027   0.0   42.9   7.0     274   41737   0.0   42.9   7.0     301   39647   0.0   42.9   7.0					
197         46330         0.0         37.5         8.0           198         42471         0.0         37.5         8.0           200         41173         0.0         42.9         7.0           202         42479         0.0         50.0         8.0           216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0           218         46015         0.0         62.5         8.0           225         45301         0.0         37.5         8.0           228         45303         0.0         37.5         8.0           250         41033         0.0         57.1         7.0           260         41035         0.0         57.1         7.0           268         41027         0.0         42.9         7.0           274         41737         0.0         42.9         7.0           301         39647         0.0         42.9         7.0			0.0		
198         42471         0.0         37.5         8.0           200         41173         0.0         42.9         7.0           202         42479         0.0         50.0         8.0           216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0           218         46015         0.0         62.5         8.0           225         45301         0.0         37.5         8.0           228         45303         0.0         37.5         8.0           250         41033         0.0         57.1         7.0           260         41025         0.0         57.1         7.0           268         41027         0.0         42.9         7.0           274         41737         0.0         42.9         7.0           301         39647         0.0         42.9         7.0					
200         41173         0.0         42.9         7.0           202         42479         0.0         50.0         8.0           216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0           218         46015         0.0         62.5         8.0           225         45301         0.0         37.5         8.0           228         45303         0.0         37.5         8.0           250         41033         0.0         57.1         7.0           260         41027         0.0         42.9         7.0           264         41027         0.0         42.9         7.0           301         39647         0.0         42.9         7.0           301         39647         0.0         42.9         7.0					
202         42479         0.0         50.0         8.0           216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0           218         46015         0.0         62.5         8.0           225         45301         0.0         37.5         8.0           228         45303         0.0         37.5         8.0           250         41033         0.0         57.1         7.0           260         41035         0.0         57.1         7.0           268         41027         0.0         42.9         7.0           274         41737         0.0         42.9         7.0           301         39647         0.0         42.9         7.0			0.0		
216         39621         0.0         42.9         7.0           217         46007         0.0         50.0         8.0           218         46015         0.0         62.5         8.0           225         45301         0.0         37.5         8.0           228         45303         0.0         37.5         8.0           250         41033         0.0         57.1         7.0           260         41025         0.0         57.1         7.0           268         41027         0.0         42.9         7.0           274         41737         0.0         42.9         7.0           301         39647         0.0         42.9         7.0	200	41173	0.0	42.9	7.0
217         46007         0.0         50.0         8.0           218         46015         0.0         62.5         8.0           225         48301         0.0         37.5         8.0           228         48303         0.0         37.5         8.0           250         41033         0.0         57.1         7.0           260         41025         0.0         57.1         7.0           268         41027         0.0         42.9         7.0           274         41737         0.0         42.9         7.0           301         39647         0.0         42.9         7.0	202	42479	0.0	50.0	8.0
218         46015         0.0         62.5         8.0           225         45301         0.0         37.5         8.0           228         45303         0.0         37.5         8.0           250         41033         0.0         57.1         7.0           260         41035         0.0         57.1         7.0           268         41027         0.0         42.9         7.0           274         41737         0.0         42.9         7.0           301         39647         0.0         42.9         7.0	216	39621	0.0	42.9	7.0
218         46013         0.0         62.3         6.0           225         48301         0.0         37.5         8.0           228         48303         0.0         37.5         8.0           250         41033         0.0         57.1         7.0           260         41027         0.0         57.1         7.0           268         41027         0.0         42.9         7.0           301         39647         0.0         42.9         7.0           301         39647         0.0         42.9         7.0	217	46007	0.0		8.0
228         45303         0.0         37.5         8.0           250         41033         0.0         57.1         7.0           260         41035         0.0         57.1         7.0           268         41027         0.0         42.9         7.0           274         41737         0.0         42.9         7.0           301         39647         0.0         42.9         7.0	218		0.0	02.3	8.0
250         41033         0.0         57.1         7.0           260         41035         0.0         57.1         7.0           268         41027         0.0         42.9         7.0           274         41737         0.0         42.9         7.0           301         39647         0.0         42.9         7.0	225	45301	0.0	37.5	8.0
260         41035         0.0         57.1         7.0           268         41027         0.0         42.9         7.0           274         41737         0.0         42.9         7.0           301         39647         0.0         42.9         7.0	228	45303	0.0	37,5	8.0
268     41027     0.0     42.9     7.0       274     41737     0.0     42.9     7.0       301     39647     0.0     42.9     7.0	250	41033	0.0		7.0
274         41737         0.0         42.9         7.0           301         39647         0.0         42.9         7.0	260	41035	0.0	57.1	7.0
301 39647 0.0 42.9 7.0	268	41027	0.0	42.9	7.0
	274	41737	0.0	42.9	7.0
307 38943 0.0 42.9 7.0	301	39647	0.0	42.9	7.0
	307	38943	0.0	42.9	7.0

Table 8

Labi	e 8			
SEQ ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< th=""><th>T/N Colon Num Ratios</th></halfx<>	T/N Colon Num Ratios
309	38939	0.0	42.9	7.0
315	44939	0.0	37.5	8.0
324	42827	0.0	37.5	8.0
326	38231	0.0	42.9	7.0
334	42819	0.0	37.5	8.0
352	43521	0.0	62.5	8.0
358	45633	0.0	50.0	8.0
364	44931	0.0	50.0	8.0
365	45635	0.0	50.0	8.0
366	46345	0.0	37.5	8.0
390	44947	0.0	50.0	8.0
391	44247	0.0	50.0	8.0
403	43 50 1	0.0	37.5	8.0
406	43489	0.0	50.0	8.0
407	44951	0.0	37.5	8.0
413	41755	0.0	42.9	7.0
420	43541	0.0	37.5	8.0
424	44953	0.0	50.0	8.0
426	46365	0.0	62.5	8.0
432	44909	0.0	50.0	8.0
435	38210	0.0	42.9	7.0
443	38928	0.0	42.9	7.0
444	44911	0.0	50.0	8.0
446	46361	0.0	50.0	8.0
450	39632	0.0	42.9	7.0
452	39620	0.0	42.9	7.0
455	46363	0.0	62.5	8.0
458	41736	0.0	57.1	7.0
464	38944	0.0	42.9	7.0
467	45605	0.0	62.5	8.0
468	45609	0.0	100.0	8.0
471	38228	0.0	57.1	7.0
472	41740	0.0	42.9	7.0
476	41032	0.0	42.9	7.0
480	39638	0.0	57.1	7.0
482	41760	0.0	42.9	7.0
490	41754	0.0	71.4	7.0
496	39980	0.0	57.1	7.0
497	46315	0.0	37.5	8.0
507	40674	0.0	42.9	7.0
509	38566	0.0	57.1	7.0
519	38590	0.0	42.9	· 7.0
539	42813	0.0	37.5	8.0

Table 8

1 abi	<del></del>			
SEQ ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< th=""><th>T/N Colon Num Ratios</th></halfx<>	T/N Colon Num Ratios
554	43515	0.0	50.0	8.0
558	41400	0.0	42.9	7.0
560	40702	0.0	42.9	7.0
563	40000	0.0	42.9	7.0
573	38185	0.0	42.9	7.0
582	39587	0.0	42.9	7.0
587	44925	0.0	50.0	8.0
592	39597	0.0	57.1	7.0
593	39593	0.0	42,9	7.0
603	38893	0.0	42.9	7.0
606	42842	0.0	62.5	8.0
607	43540	0.0	50.0	8.0
611	42840	0.0	50.0	8.0
614	43548	0.0	37.5	8.0
617	43538	0.0	50.0	8.0
618	46340	0,0	37.5	8.0
644	39586	0,0	42.9	7.0
651	45656	0.0	37.5	8.0
654	44254	0.0	50.0	8.0
655	45652	0.0	37.5	8.0
666	46285	0.0	37.5	8.0
667	40290	0.0	42.9	7.0
668	40304	0.0	42.9	7.0
680	39592	0.0	42.9	7.0
682	44950	0.0	37.5	8.0
691	45571	0.0	37.5	8.0
702	45654	0.0	37.5	8.0
703	45660	0.0	37.5	8.0
705	40292	0.0	42.9	7.0
711	40294	0.0	42.9	7.0
722	46364	0.0	37.5	8.0
724	38892	0.0	42.9	7.0
733	40998	0.0	57.1	7.0
736	40996	0.0	57.1	7.0
738	41712	0.0	42.9	7.0
757	38196	0.0	42.9	7.0
773	44881	0.0	37.5	8.0
776	39610	0.0	42.9	7.0
790	41016	0.0	42.9	7.0
793	39942	0.0	42.9	7.0
795	41718	0.0	42.9	7.0
798	39938	0.0	42.9	7.0
801	46289	0.0	37.5	8.0

Table 8

Tabl	e 8			
SEO ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< th=""><th>T/N Colon Num Ratios</th></halfx<>	T/N Colon Num Ratios
804	41024	0.0	42.9	7.0
807	38536	0.0	71.4	7.0
810	39948	0.0	42.9	7.0
816	39236	0,0	71.4	7.0
818	38540	0,0	42,9	7.0
820	41720	0.0	42.9	7.0
821	41728	0.0	42.9	7,0
831	46293	0.0	37.5	8.0
843	41358	0.0	71.4	7.0
846	39954	0.0	57.1	7.0
850	41360	0.0	42.9	7.0
864	38550	0.0	42,9	7.0
866	38409	0.0	31.7	41.0
868	40652	0.0	42.9	7.0
881	42070	0.0	57.1	7.0
883	42072	0.0	57.1	7.0
884	42074	0.0	42.9	7.0
886	40658	0.0	42.9	7.0
889	41372	0.0	42.9	7.0
-895	40670	0.0	42.9	7.0
905	38147	0.0	42.9	7.0
915	39563	0.0	42.9	7.0
916	38863	0.0	42.9	7.0
918	38859	0.0	42.9	7.0
937	40346	0.0	42.9	7.0
941	41046	0.0	42.9	7.0
945	45605	0.0	62.5	8.0
946	40326	0.0	71.4	7.0
948	40328	0,0	42.9	7.0
950	41032	0.0	42.9	7.0
955	40342	0.0	42.9	7.0
960	41742	0.0	42.9	7.0
962	41056	0.0	42.9	7.0
972 974	43215	0.0	50.0	8.0
974	43203 42497	0.0	37.5	8.0
973		0.0	37.5	8.0
984	42505 43209	0.0	62.5 50.0	8.0 8.0
985				
986	38431 24379	0.0	57.1 36.6	7.0
989	43909	0.0	100.0	8.0
991	41667	0.0	42.9	7.0
992	40985	0.0	42.9	7.0
	70,00	, 0.0	74.7	7.0

Table 8

Labi	e 8			
SEQ ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< th=""><th>T/N Colon Num Ratios</th></halfx<>	T/N Colon Num Ratios
996	38873	0.0	42.9	7.0
998	38875	0.0	42,9	7.0
999	40977	0.0	42.9	7.0
1001	38169	0.0	42.9	7.0
1005	40987	0.0	42.9	7.0
1006	40261	0.0	42.9	7.0
1010	39809	0.0	42.9	7.0
1015	40973	0.0	42.9	7.0
1016	39579	0.0	42.9	7.0
1018	40965	0.0	42.9	7.0
1024	40263	0.0	42.9	7.0
1026	39811	0.0	57.1	7.0
1028	40513	0.0	57.1	7.0
1031	39821	0.0	42.9	7.0
1032	38871	0,0	42.9	7.0
1038	38175	0.0	42.9	7.0
1053	40267	0.0	42.9	7.0
1054	40273	0.0	42.9	7.0
1057	40525	0.0	42.9	7.0
1064	41685	0.0	42.9	7.0
1066	40991	0.0	42.9	7.0
1067	41217	0.0	71.4	7.0
1072	39907	0.0	57.1	7.0
1076	41221	0,0	42.9	7.0
1085	42027	0.0	42.9	7.0
1106	41227	0.0	42.9	7.0
1112	41923	0.0	71.4	7.0
1114	41223	0.0	42.9	7.0
1124	38503	0.0	42.9	7.0
1130	41933	0.0	42.9	7.0
1132	40623	0.0	42.9	7.0
1135	38527	0.0	42.9	7.0
1138	39905	0.0	42.9	7.0
1141	40613	0.0	42,9	7.0
1142	40615	0.0	42,9	7.0
1146	39925	0.0	42.9	7.0
1147	41333	0.0	42.9	7.0
1152	40627	0.0	42.9	7.0
1153	41339	0.0	42.9	7.0
1157	39933	0.0	42.9	7.0
1159	40629	0.0	42.9	7.0
1166	42045	0.0	42.9	7.0
1167	39921	0.0	42.9	7.0

Table 8

Laur	-			
SEQ ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< th=""><th>T/N Colon Num Ratios</th></halfx<>	T/N Colon Num Ratios
1170	40637	0.0	71.4	7.0
1181	42035	0.0	42.9	7.0
1186	43931	0.0	37.5	8.0
1189	46029	0.0	37.5	8.0
1193	42523	0.0	37.5	8.0
1199	39829	0.0	42.9	7,0
1203	43923	0.0	62.5	8.0
1206	43229	0.0	50.0	8.0
1208	44629	0.0	37.5	8.0
1220	43219	0.0	. 50.0	8.0
1222	39835	0.0	100.0	7.0
1231	40529	0.0	100.0	7.0
1234	43921	0.0	37.5	8.0
1238	45319	0.0	50.0	8.0
1241	45313	0.0	37.5	8.0
1245	44627	0.0	37.5	8.0
1246	44631	0.0	37.5	8.0
1250	40531	0.0	42.9	7.0
1255	46035	0.0	62.5	8.0
1270	41233	0.0	85.7	7.0
1274	40537	0.0	42.9	7.0
1280	44637	0.0	37.5	8.0
1281	45335	0.0	37.5	8.0
1290	40535	0.0	57.1	7.0
1292	41241	0.0	42.9	7.0
1293	41943	0.0	42.9	7.0
1311	41947	0.0	42.9	7.0
1382	38765	0.0	57.1	7.0
1392	39467	0.0	57.1	7.0
1398	42861	0.0	62.5	8.0
1399	43559	0.0	37.5	8.0
1401	38146	0.0	37.5	8.0
1402	43553	0.0	37.5	8.0
1408	43555	0.0	42.9	7.0
1412	39463	0.0	71.4	7.0
1413	43557	0.0	42.9	7.0
1415	40175	0.0	42.9	7.0
1418	40167	0.0	42.9	7.0
1422	40260	0.0	37.5	8.0
1429	44965	0,0	37.5	8.0
1430	44969	0.0	42.9	7.0
1432	44967	0.0	42.9	7.0
1442	40165	0.0	42.9	7.0

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Table 8

Tabl	e 8			
SEO ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< th=""><th>T/N Colon Num Ratios</th></halfx<>	T/N Colon Num Ratios
1446	44265	0.0	42.9	7.0
1448	38162	0.0	37.5	8.0
1450	41678	0.0	37.5	8.0
1452	40974	0.0	37.5	8,0
1454	41674	0.0	37,5	8,0
1458	46379	0.0	37.5	8.0
1463	41670	0.0	37.5	8.0
1467	42871	0.0	50.0	8,0
1472	38172	0.0	37.5	8,0
1474	44273	0.0	50.0	8.0
1475	44277	0.0	50.0	8.0
1476	43569	0.0	37.5	8.0
1483	38872	0.0	50,0	8.0
1486	43577	0.0	50.0	8.0
1492	39576	0.0	57.1	7.0
1493	44977	0.0	50.0	8.0
1501	39580	0.0	62.5	8.0
1502	45689	0.0	37.5	8.0
1503	44985	0.0	50,0	8.0
1504	45681	0.0	75.0	8.0
1507	39578	0.0	57.1	7.0
1508	40984	0.0	50.0	8.0
1510	39584	0.0	42.9	7.0
1512	40990	0.0	37.5	8.0
1514	46391	0.0	37.5	8.0
1516	41682	0.0	42.9	7.0
1526	38769	0.0	42.9	7.0
1530	44612	0.0	37,5	8.0
1532	44622	0.0	37,5	8.0
1548	39473	0.0	57.1	7.0
1550	42281	0.0	42,9	7.0
1553	45320	0.0	37,5	8.0
1554	39479	0.0	42.9	7.0
1560	42287	0.0	42.9	7.0
1561	45314	0.0	37,5	8.0
1562	45326	0.0	37.5	8.0
1567	42273	0.0	42.9	7.0
1568	43210	0.0	37.5	8.0
1573	43910	0.0	37,5	8.0
1575	42279	0,0	42.9	7.0
1584	46034	0.0	37.5	8.0
1585	43934	0.0	50.0	8.0
1586	43936	0.0	50,0	8.0

#### PCT/US01/25840

Table 8

Table	e 8			
SEQ ID NO	SpotID	T/N Colon >2x	T/N Colon <a href="https://www.news.com">https://www.news.com</a>	T/N Colon Num Ratios
1587	44632	0.0	50.0	8.0
1588	43222	0.0	50.0	8.0
1589	40187	0.0	42.9	7.0
1590	44626	0.0	50.0	8.0
1597	44640	0.0	50.0	8.0
1599	43232		37.5	
	43232	0.0		8.0
1601 1603		0.0	37.5	8.0
	44628	0.0	37.5	8.0
1609	44638	0.0	37.5	8.0
1610	45332	0.0	50.0	8.0
1611	46042	0.0	37.5	8.0
1613	43228	0.0	37.5	8.0
1615	43932	0.0	37.5	8.0
1619	40183	0.0	57.1	7.0
1623	44260	0.0	37.5	8.0
1628	43562	0.0	62.5	8.0
1632	43564	0.0	37.5	8.0
1634	45666	0.0	50.0	8.0
1636	44968	0.0	37.5	8.0
1638	42852	0.0	37.5	8.0
1642	44974	0.0	50.0	8.0
1645	41587	0.0	42.9	7.0
1646	44266	0.0	37.5	8.0
1647	44268	0.0	37.5	8.0
1648	44962	0.0	37.5	8.0
1653	44972	0.0	37.5	8.0
1654	45668	0.0	50,0	8.0
1662	41593	0.0	42.9	7.0
1664	45676	0.0	50,0	8.0
1667	42866	0.0	62.5	8.0
1669	44274	0.0	37.5	8.0
1673	42874	0.0	50.0	8.0
1675	42876	0.0	37.5	8.0
1679	42289	0.0	57.1	7.0
1681	42880	0.0	37.5	8.0
1682	43580	0.0	50.0	8.0
1686	46384	0.0	37.5	8.0
1689	45682	0.0	37.5	8.0
1698	46396	0.0	50.0	8.0
1703	38406	0.0	57.1	7.0
1705	44282	0.0	37.5	8.0
1706	46400	0.0	37.5	8.0
1717	46388	0.0	37.5	8.0

Table 8

Tabl	e 8			
SEQ ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< th=""><th>T/N Colon Num Ratios</th></halfx<>	T/N Colon Num Ratios
1722	38416	0.0	42.9	7.0
1727	42301	0.0	42.9	7.0
1728	44978	0.0	37.5	8.0
1729	42543	0.0	37.5	8.0
1732	42535	0.0	37.5	8.0
1733	45684	0.0	50.0	8.0
1737	44990	0.0	50.0	8.0
1738	45686	0.0	62.5	8.0
1739	46390	0.0	37.5	8.0
1740	42531	0.0	50.0	8.0
1746	43243	0.0	37.5	8.0
1747	43947	0.0	50.0	8.0
1749	46055	0.0	50.0	8.0
1753	44651	0.0	37.5	8.0
1756	45347	0.0	37.5	8.0
1757	42547	0.0	37.5	8.0
1758	39816	0.0	42.9	7.0
1760	44643	0.0	50.0	8.0
1763	42555	0.0	37.5	8.0
1764	39114	0.0	57.1	7.0
1770	43945	0.0	37.5	8.0
1771	44647	0.0	37.5	8.0
1776	45359	0.0	37.5	8.0
1777	42551	0.0	50.0	8.0
1781	46049	0.0	37.5	8.0
1785	42545	0.0	37.5	8.0
1787	43261	0.0	37.5	8.0
1788	44657	0.0	50.0	8.0
1793	43249	0.0	37.5	8.0
1794	43255	0.0	37.5	8.0
1796	43959	0.0	50.0	8.0
1797	40524	0.0	42.9	7.0
1801	40526	0.0	42.9	7.0
1808	43961	0.0	50.0	8.0
1814	44661	0.0	50.0	8.0
1816	40520	0.0	42.9	7.0
1819	46075	0.0	50.0	8.0
1821	46079	0.0	50.0	8.0
1822	45375	0.0	37.5	8.0
1823	41222	0.0	42.9	7.0
1826	45367	0.0	37.5	8.0
1827	46067	0.0	37.5	8.0
1831	41224	0.0	42.9	7.0

Table 8

1 801	6.0			
SEO ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< td=""><td>T/N Colon Num Ratios</td></halfx<>	T/N Colon Num Ratios
1834	41230	0.0	57.1	7.0
1836	39204	0,0	42.9	7.0
1845	38504	0,0	42.9	7.0
1850	40612	0.0	42.9	7.0
1851	40616	0.0	42.9	7.0
1854	40614	0.0	42.9	7.0
1855	40624	0.0	42.9	7.0
1859	39912	0,0	42.9	7.0
1861	39918	0.0	42.9	7.0
1868	39906	0.0	42.9	7.0
1870	38528	0.0	42.9	7.0
1875	39226	0.0	42.9	7.0
1885	38514	0.0	42.9	7.0
1888	38522	0.0	42.9	7.0
1891	39230	0.0	42.9	7.0
1892	39922	0.0	42.9	7.0
1898	39924	0.0	42.9	7.0
1906	39936	0.0	42.9	7.0
1907	40626	0.0	42.9	7.0
1913	41240	0.0	57.1	7.0
1916	40225	0.0	42.9	7.0
1922	41641	0.0	42.9	7.0
1927	42036	0.0	42.9	7.0
1927	41938	0.0	42.9	7.0
1929	40235	0.0	42.9	7.0
1932	38117	0.0	42.9	7.0
1933	40929	0.0	42.9	7.0
1944	41952	0.0	42.9	7.0
1946	39527	0.0	57.1	7.0
1950	39533	0.0	42.9	7.0
1954	41944	0.0	42.9	7.0
1957	42046	0.0	42.9	7.0
1963	41342	0.0	42.9	7.0
1964	39535			7.0
1964	40544	0.0	42.9 42.9	7.0
1970	38821	0.0	42.9	7.0
1970	40231	0.0	42.9	7.0
1971	41647	0.0	42.9	7.0
1972	41047	0.0		7.0
1973	38823	0.0	42.9	7.0
1977	40943	0.0	42.9	7.0
1988	38831	0.0	42.9	7.0
1988	38127		42.9	
1990	3812/	0.0	42.9	7.0

Table 8

Tabl	e 8			
SEO ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< th=""><th>T/N Colon Num Ratios</th></halfx<>	T/N Colon Num Ratios
1992	42044	0.0	42.9	7.0
2007	40945	0.0	42.9	7.0
2008	40953	0.0	42.9	7.0
2013	38135	0.0	42.9	7.0
2015	40959	0.0	42.9	7.0
2022	40249	0.0	42.9	7.0
2034	39551	0.0	42.9	7.0
2036	40949	0.0	42.9	7.0
2038	41651	0.0	42.9	7.0
2042	38143	0.0	42.9	7.0
2043	38835	0.0	42.9	7.0
2045	38843	0.0	42.9	7.0
2060	41987	0.0	42.9	7.0
2068	40587	0.0	42.9	7.0
2072	39875	0.0	42.9	7.0
2076	40589	0.0	42.9	7.0
2077	38471	0.0	42.9	7.0
2079	38483	0.0	42.9	7.0
2081	41283	0.0	42.9	7.0
2120	39195	0.0	42.9	7.0
2121	39891	0.0	42.9	7.0
2125	39193	0.0	42.9	7.0
2130	39199	0,0	42.9	7.0
2158	40593	0.0	42,9	7.0
2163	40603	0.0	42,9	7.0
2177	42009	0.0	42.9	7.0
2185	39526	0.0	42.9	7.0
2187	39536	0.0	42.9	7.0
2193	40942	0.0	42.9	7.0
2195	38120	0.0	42.9	7.0
2199	40238	0.0	42.9	7.0
2202	40240	0.0	42.9	7.0
2204	39522	0.0	42.9	7.0
2206	39534	0.0	42.9	7.0
2208	42011	0.0	42.9	7.0
2214	42013	0.0	42.9	7.0
2224	38132	0.0	42.9	7.0
2229	40256	0.0	42.9	7.0
2235	42343	0.0	37.5	8.0
2236	43041	0.0	37.5	8.0
2237	38144	0.0	42.9	7.0
2239	39548	0.0	57.1	7.0
2243	38842	0.0	42.9	7.0

Table 8

Tabl	e 8			
SEO ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< th=""><th>T/N Colon Num Ratios</th></halfx<>	T/N Colon Num Ratios
2247	38142	0.0	42.9	7.0
2250	38138	0.0	42.9	7.0
2257	39552	0.0	57.1	7.0
2263	39544	0.0	42.9	7.0
2270	41650	0.0	42.9	7.0
2271	40952	0.0	42.9	7.0
2272	41652	0.0	42.9	7.0
2278	38470	0.0	42.9	7.0
2279	40954	0.0	42.9	7.0
2288	43753	0.0	50.0	8.0
2290	39176	0.0	57.1	7.0
2293	39874	0.0	42.9	7.0
2294	40590	0.0	42.9	7.0
2300	40592	0.0	42.9	7.0
2304	41292	0.0	42.9	7.0
2312	39180	0.0	42.9	7.0
2314	41294	0.0	42.9	7.0
2323	39192	0.0	42.9	7.0
2326	39888	0.0	42.9	7.0
2327	40584	0.0	42.9	7.0
2328	41282	0.0	42.9	7.0
2329	41990	0.0	42.9	7.0
2336	39184	0.0	42.9	7.0
2338	44459	0.0	62.5	8.0
2340	40586	0.0	42.9	7.0
2341	41992	0.0	42.9	7.0
2342	44457	0.0	37.5	8.0
2342	40580	0.0	42.9	7.0
2346	39186	0.0	85.7	7.0
2347	43747	0.0	50.0	8.0
2349	41288	0.0	42.9	7.0
2350	41288	0.0	42.9	7.0
2352	38494	0.0	71.4	7.0
2352				7.0
	39188	0.0	85.7	
2354	41996 40608	0.0	42.9 42.9	7.0
2364	45165	0.0	50.0	8.0
2364	41312	0.0	50.0 42.9	7.0
2368	39198	0.0	42.9	7.0
2368	41306	0.0	42.9	7.0
2370	39904	0.0	57.1	7.0
		0.0		
2374	45163	0.0	37.5	8.0
2376	42002	0.0	42.9	7.0

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Table 8

Tabl	e 8			
SEQ ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< th=""><th>T/N Colon Num Ratios</th></halfx<>	T/N Colon Num Ratios
2380	41310	0.0	42.9	7.0
2383	39200	0.0	42.9	7.0
2386	42899	0.0	50.0	8.0
2388	45869	0.0	37.5	8.0
2390	42901	0.0	37.5	8.0
2393	45709	0.0	62.5	8.0
2400	42893	0.0	50.0	8.0
2404	43585	0.0	37.5	8.0
2408	43599	0.0	50.0	8.0
2412	43587	0.0	37.5	8.0
2413	44301	0.0	50.0	8.0
2414	46411	0.0	37.5	8.0
2415	42909	0.0	62.5	8.0
2422	42895	0.0	37.5	8.0
2423	44995	0.0	37.5	8.0
2423	42911	0.0	50.0	8.0
2424	45865	0.0	37.5	8.0
2429	42885	0.0	37.5	8.0
2432	42887	0.0	50.0	8.0
2432	45705	0.0	37.5	8.0
2439	42012	0.0	42.9	7.0
2439	43593		37.5	8.0
2442		0.0	37.5	8.0
2448	42905	0.0		8.0
2456	44303	0.0	50.0	
	45697	0.0	50.0 62.5	8.0
2468	44305	0.0		8.0
2469 2471	43609 45009	0.0	37.5	8.0
			50.0	8.0
2475 2476	44317 45719	0.0	50,0	8.0
			37.5	8.0
2477	43761	0.0	50.0	8.0
2478	46431	0.0	50.0	8.0
2480	44311	0.0	62.5	8.0
2483	45017	0.0	50.0	8.0
2484	46421	0.0	50.0	8.0
2485	44313	0.0	50.0	8.0
2487	45019	0.0	37,5	8.0
2489	46417	0.0	37.5	8.0
2491	46423	0.0	37.5	8.0
2496	45713	0.0	37.5	8.0
2498	46425	0.0	37.5	8.0
2500	45013	0.0	37.5	8.0
2502	44319	0.0	37.5	8.0

Table 8

			r	
SEQ ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< td=""><td>T/N Colon Num Ratios</td></halfx<>	T/N Colon Num Ratios
2503	45723	0.0	37.5	8.0
2504	46427	0.0	37.5	8.0
2505	42534	0.0	37.5	8.0
2511	43938	0.0	37.5	8.0
2516	43950	0.0	37.5	8.0
2517	45360	0.0	37.5	8.0
2518	43944	0.0	37.5	8.0
2524	45348	0.0	37.5	8.0
2527	46052	0.0	37.5	8.0
2532	43946	0.0	37.5	8.0
2534	43236	0.0	37.5	8.0
2539	44650	0.0	37.5	8.0
2541	43242	0.0	50.0	8.0
2544	43244	0.0	50.0	8.0
2549	46056	0.0	50.0	8.0
2550	42556	0.0	37.5	8.0
2551	43262	0.0	50.0	8.0
2555	45169	0.0	37.5	8.0
2560	42550	0.0	37.5	8.0
2563	42552	0.0	37.5	8.0
2564	43966	0.0	50.0	8.0
2581	43968	0.0	37.5	8.0
2582	42554	0.0	50.0	8.0
2591	44660	0.0	37.5	8.0
2592	45362	0.0	37.5	8.0
2596	45376	0.0	62.5	8.0
2601	45364	0.0	37.5	8.0
2604	45374	0.0	37.5	8.0
2605	46066	0.0	37.5	8.0
2608	45368	0.0	37.5	8.0
2609	46078	0.0	50.0	8.0
2613	44668	0.0	50.0	8.0
2614	45370	0.0	37.5	8.0
2619	43592	0.0	37.5	8.0
2620	45875	0.0	37.5	8.0
2621	46068	0.0	75.0	8.0
2623	42882	0.0	37.5	8.0
2629	43588	0.0	37.5	8.0
2638	44296	0.0	37.5	8.0
2639	43401	0.0	50.0	8.0
2641	44298	0.0	37.5	8.0
2642	45710	0.0	50.0	8.0
2646	45008	0.0	50,0	8.0

Table 8

Labi	e o			
SEO ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< td=""><td>T/N Colon Num Ratios</td></halfx<>	T/N Colon Num Ratios
2647	42910	0.0	62.5	8.0
2648	43596	0,0	75.0	8,0
2649	45706	0,0	37.5	8,0
2655	46410	0.0	37.5	8,0 .
2658	44994	0.0	50.0	8.0
2662	45708	0.0	50.0	8.0
2666	46416	0.0	37.5	8.0
2667	42912	0.0	37.5	8.0
2671	42902	0.0	37.5	8.0
2673	44300	0.0	37.5	8.0
2676	42904	0.0	50.0	8.0
2677	43598	0.0	50.0	8.0
2686	42906	0.0	50.0	8.0
2688	42695	0.0	37.5	8.0
2689	46404	0.0	37.5	8.0
2690	42898	0.0	50.0	8.0
2694	45885	0.0	37.5	8.0
2703	42908	0.0	37.5	8.0
2704	44292	0.0	37.5	8.0
2710	46412	0.0	37.5	8.0
2711	44306	0.0	37.5	8.0
2713	43614	0.0	37.5	8.0
2715	43616	0.0	37.5	8.0
2718	43612	0.0	50.0	8.0
2726	43606	0.0	87.5	8.0
2727	43610	0.0	50.0	8.0
2729	43602	0.0	37.5	8.0
2732	45714	0.0	37.5	8.0
2733	43604	0.0	50.0	8.0
2739	45018	0.0	50.0	8.0
2755	45718	0.0	37.5	8.0
2812	46418	0.0	37.5	8.0
2885	46215	0.0	37.5	8.0
2966	45529	0.0	37.5	8.0
2983	46229	0.0	37.5	8.0
2984	45533	0.0	50.0	8.0
3006	43756	0.0	50.0	8.0
3011	43758	0.0	62.5	8.0
3082	46222	0.0	37.5	8.0
3093	46212	0.0	37.5	8.0
3129	42718	0.0	37.5	8.0
3145	42710	0.0	37.5	8.0
3176	43424	0.0	37.5	8.0

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Table 8

Tabl	e 8			
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SEQ ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< td=""><td>T/N Colon Num Ratios</td></halfx<>	T/N Colon Num Ratios
3193	44822	0.0	37.5	8.0
3211	44818	0.0	50.0	8.0
3245	45860	0.0	50.0	8.0
3257	45858	0.0	37.5	8.0
3268	42356	0.0	37.5	8.0
3269	42364	0.0	37.5	8.0
3278	45862	0.0	37.5	8.0
3296	43064	0.0	37.5	8.0
3336	45170	0.0	37.5	8.0
3339	43768	0.0	50.0	8.0
3356	43776	0.0	50.0	8.0
3359	45176	0.0	50.0	8.0
3388	42698	0.0	50.0	8.0
3393	45184	0.0	37.5	8.0
3405	45878	0.0	37.5	8.0
3412	45884	0.0	37.5	8.0
3438	46020	0.0	37.5	8.0
3439	46032	0.0	50.0	8.0
3442	46026	0.0	37.5	8.0
3443	42516	0.0	50.0	8.0
3457	44117	0.0	50.0	8.0
3463	42719	0.0	50.0	8.0
3514	43423	0.0	50.0	8.0
3575	46233	0.0	37.5	8.0
3595	43054	0.0	50.0	8.0
3596	42352	0.0	37.5	8.0
3605	43746	0.0	50.0	8.0
3613	42366	0.0	50.0	8.0
3650	40101	0.0	42.9	7.0
3656	39407	0.0	42.9	7.0
3698	38695	0.0	42.9	7.0
3702	40107	0.0	42.9	7.0
3711	39401	0.0	42.9	7.0
3716	39405	0.0	42.9	7.0
3723	38697	0.0	42.9	7.0
3836	42213	0.0	42.9	7.0
3870	38717	0.0	42.9	7.0
3881	38719	0.0	42.9	7.0
3919	38707	0.0	42.9	7.0
3934	38713	0.0	42.9	7.0
3955	39419	0.0	42.9	7.0
4028	42274	0.0	42.9	7.0
4039	38772	0.0	42.9	7.0

Table 8

Tabl	e 8			
SEQ ID NO	SpotID	T/N Colon >2x	T/N Colon <a href="halfx">halfx</a>	T/N Colon Num Ratios
4041	42286	0.0	42.9	7.0
4045	38770	0.0	42.9	7.0
4055	42282	0.0	42.9	7.0
4059	42284	0.0	42.9	7.0
4097	38774	0.0	42.9	7.0
4124	40313	0.0	42.9	7.0
4131	45625	0.0	50.0	8.0
4137	41005	0.0	42.9	7.0
4140	38203	0.0	42.9	7.0
4145	46325	0.0	37.5	8.0
4146	39611	0.0	42.9	7.0
4147	40309	0.0	57.1	7.0
4153	45619	0.0	37.5	8.0
4163	41697	0.0	42.9	7.0
4166	38899	0.0	42.9	7.0
4170	38903	0.0	42.9	7.0
4171	41003	0.0	42.9	7.0
4173	40995	0.0	42.9	7.0
4175	46321	0.0	37.5	8.0
4177	41017	0.0	42.9	7.0
4182	42474	0.0	50.0	8.0
4195	42478	0.0	37.5	8.0
4197	41015	0.0	42.9	7.0
4212	41713	0.0	42.9	7.0
4214	42480	0.0	37.5	8.0
4217	43886	0.0	37.5	8.0
4228	43888	0.0	37.5	8.0
4240	44586	0.0	37.5	8.0
4252	43188	0.0	37.5	8.0
4254	45304	0.0	62.5	8,0
4263	45996	0.0	37.5	8.0
4275	46016.	0.0	37.5	8.0
4277	43198	0.0	37.5	8.0
4279	44606	0.0	50.0	8.0
4282	42496	0.0	50.0	8.0
4285	43900	0.0	37.5	8.0
4286	44608	0.0	37.5	8,0
4288	43902	0.0	75.0	8.0
4290	46006	0.0	37.5	8.0
4293	43192	0.0	37.5	8.0
4296	42490	0.0	37.5	8.0
4299	43896	0.0	37.5	8.0
4305	42492	0.0	37.5	8.0

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Table 8

4307         46008         0.0         50.0         8.0           4317         39941         0.0         42.9         7.0           4324         42053         0.0         42.9         7.0           4341         38541         0.0         42.9         7.0           4343         42055         0.0         42.9         7.0           4371         40647         0.0         42.9         7.0           4384         41355         0.0         42.9         7.0           4386         39261         0.0         42.9         7.0           4395         39967         0.0         42.9         7.0           4397         38559         0.0         42.9         7.0           4403         40669         0.0         42.9         7.0           4403         40669         0.0         42.9         7.0           4421         44930         0.0         37.5         8.0           4422         4201         0.0         42.9         7.0           4424         42071         0.0         42.9         7.0           4424         42071         0.0         42.9         7.0 <th></th> <th></th> <th></th> <th></th> <th></th>					
4317         39941         0.0         42.9         7.0           4324         42053         0.0         42.9         7.0           4341         38541         0.0         42.9         7.0           4343         38541         0.0         42.9         7.0           4350         39241         0.0         42.9         7.0           4371         40647         0.0         42.9         7.0           4384         41355         0.0         42.9         7.0           4386         39261         0.0         42.9         7.0           4397         38559         0.0         42.9         7.0           4400         40663         0.0         42.9         7.0           4403         40669         0.0         42.9         7.0           4403         40669         0.0         42.9         7.0           4403         43963         0.0         42.9         7.0           4421         44930         0.0         37.5         8.0           4422         45638         0.0         37.5         8.0           4422         45638         0.0         37.5         8.0 </td <td>SEQ ID NO</td> <td>SpotID</td> <td>T/N Colon &gt;2x</td> <td>T/N Colon <halfx< td=""><td>T/N Colon Num Ratios</td></halfx<></td>	SEQ ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< td=""><td>T/N Colon Num Ratios</td></halfx<>	T/N Colon Num Ratios
4324         42053         0.0         42.9         7.0           4341         38541         0.0         42.9         7.0           4343         38541         0.0         42.9         7.0           4350         39241         0.0         42.9         7.0           4371         40647         0.0         42.9         7.0           4384         41355         0.0         42.9         7.0           4386         39261         0.0         42.9         7.0           4395         39967         0.0         42.9         7.0           4400         40663         0.0         42.9         7.0           4400         40663         0.0         42.9         7.0           4403         40669         0.0         42.9         7.0           4421         44930         0.0         37.5         8.0           4422         4201         0.0         42.9         7.0           4424         42071         0.0         42.9         7.0           4424         42071         0.0         42.9         7.0           4422         41371         0.0         42.9         7.0 <td>4307</td> <td>46008</td> <td>0.0</td> <td>50.0</td> <td>8.0</td>	4307	46008	0.0	50.0	8.0
4341         38541         0.0         42.9         7.0           4343         42055         0.0         42.9         7.0           4350         39241         0.0         42.9         7.0           4371         40647         0.0         42.9         7.0           4384         41355         0.0         42.9         7.0           4386         39261         0.0         42.9         7.0           4395         39967         0.0         42.9         7.0           4397         38559         0.0         42.9         7.0           4400         40663         0.0         42.9         7.0           4403         40669         0.0         42.9         7.0           4403         40669         0.0         42.9         7.0           4421         44930         0.0         37.5         8.0           4424         42071         0.0         42.9         7.0           4422         45638         0.0         37.5         8.0           4429         41371         0.0         42.9         7.0           44428         44268         0.0         37.5         8.0     <	4317	39941	0.0	42.9	7.0
4343         42055         0.0         42.9         7.0           4350         39241         0.0         42.9         7.0           4371         40647         0.0         42.9         7.0           4384         41355         0.0         42.9         7.0           4386         39261         0.0         42.9         7.0           4397         38559         0.0         42.9         7.0           4400         40663         0.0         42.9         7.0           4400         40663         0.0         42.9         7.0           4408         39263         0.0         42.9         7.0           4421         44930         0.0         37.5         8.0           4422         45638         0.0         37.5         8.0           4422         45638         0.0         37.5         8.0           4428         44228         0.0         37.5         8.0           4429         41371         0.0         42.9         7.0           4434         45640         0.0         50.0         8.0           4434         44660         0.0         50.0         8.0 </td <td>4324</td> <td>42053</td> <td>0.0</td> <td>42.9</td> <td>7.0</td>	4324	42053	0.0	42.9	7.0
4350         39241         0.0         42.9         7.0           4371         40647         0.0         42.9         7.0           4384         41355         0.0         42.9         7.0           4386         39261         0.0         42.9         7.0           4395         39967         0.0         42.9         7.0           4397         38559         0.0         42.9         7.0           4400         40663         0.0         42.9         7.0           4403         40669         0.0         42.9         7.0           4421         44930         0.0         37.5         8.0           4421         44930         0.0         37.5         8.0           4422         42071         0.0         42.9         7.0           4422         42638         0.0         37.5         8.0           4422         41371         0.0         42.9         7.0           4429         41371         0.0         42.9         7.0           4434         45640         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0 </td <td>4341</td> <td>38541</td> <td>0.0</td> <td>42.9</td> <td>7.0</td>	4341	38541	0.0	42.9	7.0
4371         40647         0.0         42.9         7.0           4384         41355         0.0         42.9         7.0           4386         39261         0.0         42.9         7.0           4395         39967         0.0         42.9         7.0           4397         38559         0.0         42.9         7.0           4403         40669         0.0         42.9         7.0           4403         40669         0.0         42.9         7.0           4408         39263         0.0         42.9         7.0           4421         44930         0.0         37.5         8.0           4424         42071         0.0         42.9         7.0           4425         45638         0.0         37.5         8.0           4429         41371         0.0         42.9         7.0           4434         45640         0.0         50.0         8.0           4434         45660         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0           4439         42818         0.0         50.0         8.0 </td <td>4343</td> <td>42055</td> <td>0.0</td> <td>42.9</td> <td>7.0</td>	4343	42055	0.0	42.9	7.0
4384         41355         0.0         42.9         7.0           4386         39261         0.0         42.9         7.0           4395         39967         0.0         42.9         7.0           4397         38559         0.0         42.9         7.0           4400         40663         0.0         42.9         7.0           4408         49669         0.0         42.9         7.0           4421         44930         0.0         37.5         8.0           4422         42071         0.0         42.9         7.0           4422         45638         0.0         37.5         8.0           4422         4428         0.0         37.5         8.0           4429         41371         0.0         42.9         7.0           4434         45640         0.0         50.0         8.0           4434         44660         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0           4436         44171         0.0         37.5         8.0           4444         45644         0.0         50.0         8.0 <td>4350</td> <td>39241</td> <td>0.0</td> <td>42.9</td> <td>7.0</td>	4350	39241	0.0	42.9	7.0
4386         39261         0.0         42.9         7.0           4395         39967         0.0         42.9         7.0           4397         38559         0.0         42.9         7.0           4400         40663         0.0         42.9         7.0           4403         40669         0.0         42.9         7.0           4421         44930         0.0         37.5         8.0           4421         44930         0.0         37.5         8.0           4422         42071         0.0         42.9         7.0           4425         45638         0.0         37.5         8.0           4422         41371         0.0         42.9         7.0           4429         41371         0.0         42.9         7.0           4434         45640         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0           4434         45640         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0           4434         45644         0.0         50.0         8.0 </td <td>4371</td> <td>40647</td> <td>0.0</td> <td>42.9</td> <td>7.0</td>	4371	40647	0.0	42.9	7.0
4395         39967         0.0         42.9         7.0           4397         38559         0.0         42.9         7.0           4400         40663         0.0         42.9         7.0           4403         40669         0.0         42.9         7.0           4408         39263         0.0         42.9         7.0           4421         44930         0.0         37.5         8.0           4422         45638         0.0         37.5         8.0           4422         45638         0.0         37.5         8.0           4429         41371         0.0         42.9         7.0           4434         45640         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0           4436         44171         0.0         37.5         8.0           4439         42818         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0 </td <td>4384</td> <td>41355</td> <td>0.0</td> <td>42.9</td> <td>7.0</td>	4384	41355	0.0	42.9	7.0
4397         38559         0.0         42.9         7.0           4400         40663         0.0         42.9         7.0           4400         40669         0.0         42.9         7.0           4408         39263         0.0         42.9         7.0           4421         44930         0.0         37.5         8.0           4421         44930         0.0         37.5         8.0           4422         45638         0.0         37.5         8.0           4428         44228         0.0         37.5         8.0           4429         41371         0.0         42.9         7.0           4434         45640         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0           4435         44163         0.0         37.5         8.0           4439         42818         0.0         50.0         8.0           4441         45634         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4441         45644         0.0         37.5         8.0 </td <td>4386</td> <td>39261</td> <td>0.0</td> <td>42.9</td> <td>7.0</td>	4386	39261	0.0	42.9	7.0
4400         40663         0.0         42.9         7.0           4403         40669         0.0         42.9         7.0           4403         40669         0.0         42.9         7.0           4421         44930         0.0         37.5         8.0           4421         44930         0.0         37.5         8.0           4422         42671         0.0         42.9         7.0           4428         44288         0.0         37.5         8.0           4429         41371         0.0         42.9         7.0           4434         45640         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0           4436         44171         0.0         37.5         8.0           4439         42818         0.0         50.0         8.0           4440         45634         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4447         43471         0.0         37.5         8.0 </td <td>4395</td> <td>39967</td> <td>0.0</td> <td>42.9</td> <td>7.0</td>	4395	39967	0.0	42.9	7.0
4403         40669         0.0         42.9         7.0           4408         39263         0.0         42.9         7.0           4421         44930         0.0         37.5         8.0           4424         42071         0.0         42.9         7.0           4425         45638         0.0         37.5         8.0           4428         44288         0.0         37.5         8.0           4429         41371         0.0         42.9         7.0           4434         45640         0.0         50.0         8.0           4434         45640         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0           4436         44171         0.0         37.5         8.0           4439         42818         0.0         50.0         8.0           4441         45634         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4447         43471         0.0         37.5         8.0           44453         44944         0.0         37.5         8.0     <	4397	38559	0.0	42.9	7.0
4408         39263         0.0         42.9         7.0           4421         44930         0.0         37.5         8.0           4421         44930         0.0         37.5         8.0           4425         45638         0.0         37.5         8.0           4428         44228         0.0         37.5         8.0           4429         41371         0.0         42.9         7.0           4434         45640         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0           4436         44171         0.0         37.5         8.0           4439         42818         0.0         50.0         8.0           4440         45634         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4441         45644         0.0         37.5         8.0           4448         43536         0.0         62.5         8.0           4448         43536         0.0         37.5         8.0           4454         45646         0.0         37.5         8.0 </td <td>4400</td> <td>40663</td> <td>0.0</td> <td>42.9</td> <td>7.0</td>	4400	40663	0.0	42.9	7.0
4421         44930         0.0         37.5         8.0           4424         42071         0.0         42.9         7.0           4424         42071         0.0         42.9         7.0           4425         45638         0.0         37.5         8.0           4428         44228         0.0         37.5         8.0           4429         41371         0.0         42.9         7.0           4434         45640         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0           4436         44171         0.0         37.5         8.0           4439         42818         0.0         50.0         8.0           4440         45634         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4447         43471         0.0         37.5         8.0           4448         43536         0.0         62.5         8.0           4448         43536         0.0         37.5         8.0 </td <td>4403</td> <td>40669</td> <td>0.0</td> <td>42.9</td> <td>7.0</td>	4403	40669	0.0	42.9	7.0
4424         42071         0.0         42.9         7.0           4425         45638         0.0         37.5         8.0           4428         45638         0.0         37.5         8.0           4429         41371         0.0         42.9         7.0           4434         45640         0.0         50.0         8.0           4434         45640         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0           4439         42818         0.0         50.0         8.0           4440         45634         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4441         45644         0.0         37.5         8.0           4447         43471         0.0         37.5         8.0           4448         43536         0.0         62.5         8.0           44453         44944         0.0         37.5         8.0           4453         44944         0.0         37.5         8.0           4454         45646         0.0         37.5         8.0     <	4408	39263	0.0	42.9	7.0
4425         45638         0.0         37.5         8.0           4428         44228         0.0         37.5         8.0           4429         4421         0.0         37.5         8.0           4434         45640         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0           4436         44171         0.0         37.5         8.0           4439         42818         0.0         50.0         8.0           4440         45634         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4441         43671         0.0         37.5         8.0           4448         43536         0.0         62.5         8.0           4445         44564         0.0         37.5         8.0           4453         44944         0.0         37.5         8.0           4457         44238         0.0         37.5         8.0           4457         44238         0.0         37.5         8.0           4458         4936         0.0         50.0         8.0 <td>4421</td> <td>44930</td> <td>0.0</td> <td>37.5</td> <td>8.0</td>	4421	44930	0.0	37.5	8.0
4428         44228         0.0         37.5         8.0           4429         41371         0.0         42.9         7.0           4434         45640         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0           4436         44171         0.0         37.5         8.0           4439         42818         0.0         50.0         8.0           4440         45634         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4441         45644         0.0         37.5         8.0           4448         43536         0.0         62.5         8.0           44453         44944         0.0         37.5         8.0           4454         45646         0.0         37.5         8.0           4457         44238         0.0         37.5         8.0           4458         44936         0.0         50.0         8.0           4461         44161         0.0         37.5         8.0	4424	42071	0.0	42.9	7.0
4429         41371         0.0         42.9         7.0           4434         45640         0.0         50.0         8.0           4434         45640         0.0         50.0         8.0           4435         44161         0.0         37.5         8.0           4439         42818         0.0         50.0         8.0           4440         45634         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4441         43471         0.0         37.5         8.0           4448         43336         0.0         62.5         8.0           4453         44944         0.0         37.5         8.0           4457         44238         0.0         37.5         8.0           4458         44936         0.0         50.0         8.0           4458         44936         0.0         50.0         8.0           4469         44938         0.0         50.0         8.0	4425	45638	0.0	37.5	8.0
4434         45640         0.0         50.0         8.0           4435         44163         0.0         37.5         8.0           4436         44171         0.0         37.5         8.0           4439         42818         0.0         50.0         8.0           4440         45634         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4441         43641         0.0         37.5         8.0           4448         43536         0.0         62.5         8.0           4453         44944         0.0         37.5         8.0           4454         45646         0.0         37.5         8.0           4457         44238         0.0         37.5         8.0           4458         4936         0.0         50.0         8.0           4461         44161         0.0         37.5         8.0           4469         44938         0.0         50.0         8.0	4428	44228	0.0	37.5	8.0
4435         44163         0.0         37.5         8.0           4436         44171         0.0         37.5         8.0           4436         44171         0.0         37.5         8.0           4439         42818         0.0         50.0         8.0           4440         45634         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4447         43471         0.0         37.5         8.0           4448         43536         0.0         62.5         8.0           4453         44944         0.0         37.5         8.0           4454         45646         0.0         37.5         8.0           4457         44238         0.0         37.5         8.0           4458         44936         0.0         50.0         8.0           4461         44161         0.0         37.5         8.0           4469         44938         0.0         50.0         8.0	4429	41371	0.0	42.9	7.0
4436         44171         0.0         37.5         8.0           4439         42818         0.0         50.0         8.0           4440         45634         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4441         43471         0.0         37.5         8.0           4448         43336         0.0         62.5         8.0           4453         44944         0.0         37.5         8.0           4454         45646         0.0         37.5         8.0           4457         4238         0.0         37.5         8.0           4458         44936         0.0         50.0         8.0           4461         44161         0.0         37.5         8.0           4469         44938         0.0         50.0         8.0	4434	45640	0.0	50.0	8.0
4439         42818         0.0         50.0         8.0           4440         45634         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4447         43471         0.0         37.5         8.0           4448         43536         0.0         62.5         8.0           4453         44944         0.0         37.5         8.0           4457         45646         0.0         37.5         8.0           4457         44238         0.0         37.5         8.0           4458         49936         0.0         50.0         8.0           4461         44161         0.0         37.5         8.0           4469         44938         0.0         50.0         8.0	4435	44163	0.0	37.5	8.0
4440         45634         0.0         50.0         8.0           4441         45644         0.0         50.0         8.0           4441         43644         0.0         50.0         8.0           4447         43471         0.0         37.5         8.0           4448         43536         0.0         62.5         8.0           4453         44944         0.0         37.5         8.0           4454         45646         0.0         37.5         8.0           4457         44238         0.0         37.5         8.0           4458         44936         0.0         50.0         8.0           4461         44161         0.0         37.5         8.0           4469         44938         0.0         50.0         8.0	4436	44171	0.0	37.5	8.0
4441         45644         0.0         50.0         8.0           4447         43471         0.0         37.5         8.0           4448         43536         0.0         62.5         8.0           44453         44944         0.0         37.5         8.0           4454         45646         0.0         37.5         8.0           4457         44238         0.0         37.5         8.0           4458         44936         0.0         50.0         8.0           4461         44161         0.0         37.5         8.0           4469         44938         0.0         50.0         8.0	4439	42818	0.0	50.0	8.0
4447     43471     0.0     37.5     8.0       4448     43536     0.0     62.5     8.0       4453     44944     0.0     37.5     8.0       4454     45646     0.0     37.5     8.0       4457     44238     0.0     37.5     8.0       4458     44936     0.0     50.0     8.0       4461     44161     0.0     37.5     8.0       4469     44938     0.0     50.0     8.0	4440	45634	0.0	50.0	8.0
4448         43536         0.0         62.5         8.0           4453         44944         0.0         37.5         8.0           4454         45646         0.0         37.5         8.0           4457         44238         0.0         37.5         8.0           4458         44936         0.0         50.0         8.0           4461         44161         0.0         37.5         8.0           4469         44938         0.0         50.0         8.0	4441	45644	0.0	50.0	8.0
4453     44944     0.0     37.5     8.0       4454     45646     0.0     37.5     8.0       4457     4228     0.0     37.5     8.0       4458     44936     0.0     50.0     8.0       4461     44161     0.0     37.5     8.0       4469     44938     0.0     50.0     8.0	4447	43471	0.0	37.5	8.0
4454     45646     0.0     37.5     8.0       4457     44238     0.0     37.5     8.0       4458     44936     0.0     50.0     8.0       4461     44161     0.0     37.5     8.0       4469     44938     0.0     50.0     8.0	4448	43536	0.0	62.5	8.0
4457     44238     0.0     37.5     8.0       4458     44936     0.0     50.0     8.0       4461     44161     0.0     37.5     8.0       4469     44938     0.0     50.0     8.0	4453	44944	0.0	37.5	8,0
4458     44936     0.0     50.0     8.0       4461     44161     0.0     37.5     8.0       4469     44938     0.0     50.0     8.0	4454	45646	0.0	37.5	8.0
4461         44161         0.0         37.5         8.0           4469         44938         0.0         50.0         8.0	4457	44238	0.0	37.5	8.0
4469 44938 0.0 50.0 8.0	4458	44936	0.0	50.0	8.0
	4461	44161	0.0	37.5	8.0
4470 45636 0.0 50.0 2.0	4469	44938	0.0		8.0
	4470	45636	0.0	50.0	8.0
4477 44804 0.0 37.5 8.0	4477	44804	0.0	37.5	8.0
4483 44100 0.0 50.0 8.0	4483		0.0	50.0	8.0
4544 46230 0.0 37.5 8.0	4544	46230	0.0	37.5	8.0
4547 45532 0.0 37.5 8.0	4547	45532	0.0	37.5	8.0
4557 45526 0.0 50.0 8.0	4557	45526	0.0	50.0	8.0
4573 45536 0.0 37.5 8.0	4573	45536	0.0	37.5	8.0
4593 41535 0.0 42.9 7.0	4593	41535	0.0	42.9	7.0
4594 40123 0.0 57.1 7.0	4594	40123	0.0	57.1	7.0
1600 11505 00 100	4600	41525	0.0	42.9	7.0

Table 8

Tabl	e 8			
SEQ ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< th=""><th>TALCIA Non Dation</th></halfx<>	TALCIA Non Dation
4611	40817	0.0	42.9	T/N Colon Num Ratios 7.0
4620	40821	0.0	42.9	7.0
4629	41529	0.0	42.9	7.0
4632	40825	0.0	42.9	7.0
4645	41527	0.0	42.9	7.0
4696	41527	0.0	42.9	7.0
4718	40823	0.0	42.9	7.0
4727	38758	0.0	57.1	7.0
4729	42229	0.0	42.9	7.0
4743	38764	0.0	42.9	7.0
4754	42235	0.0	42.9	7.0
4768	42239	0.0	42.9	7.0
4835	39472	0.0	42.9	7.0
4922	40868	0.0	57.1	7.0
4938	40866	0.0	42.9	7.0
4950	41576	0.0	42.9	7.0
4978	40870	0.0	42.9	7.0
5065	39488	0.0	42.9	7.0
5117	40888	0.0	42.9	7.0
5140	40886	0.0	42.9	7.0
5150	40890	0.0	42.9	7.0
5165	41588	0.0	42.9	7.0
5186	41596	0.0	42.9	7.0
5228	42290	0.0	42.9	7.0
5252	43118	0.0	50.0	8.0
5300	43114	0.0	62.5	8.0
5341	45220	0.0	37,5	8.0
5360	44518	0.0	37.5	8.0
5539	43120	0.0	37.5	8.0
5559	43812	0.0	50.0	8.0
5596	43810	0.0	50.0	8.0
5643	45224	0.0	50.0	8.0
5649	45226	0.0	37.5	8.0
5665	45922	0.0	37.5	8.0
5671	43265	0.0	37.5	8.0
5701	42573	0.0	37.5	8.0
5724	45232	0.0	37.5	8.0
5752	41161	0.0	71.4	7.0
5763	41163	0.0	42.9	7.0
5812	44591	0.0	50,0	8.0
5815	43189	0.0	37.5	8.0
5817	45293	0.0	37.5	8.0
5818	42487	0.0	37.5	8.0

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Table 8

Labi				
SEQ ID NO	SpotID	T/N Colon >2x	T/N Colon <halfx< td=""><td>T/N Colon Num Ratios</td></halfx<>	T/N Colon Num Ratios
5821	43191	0.0	37.5	8.0
5825	38917	0.0	42.9	7.0
5829	38913	0.0	42,9	7.0
5836	41875	0.0	42.9	7.0
5837	45987	0.0	37.5	8.0
5847	45289	0.0	37.5	8.0
5848	45989	0.0	50.0	8.0
5979	44537	0.0	50.0	8.0
80	44681	12.5	37.5	8.0
86	43981	12.5	50.0	8.0
78	44675	37.5	0.0	8.0
104	42428	37.5	0.0	8.0
3248	45866	37.5	0.0	8.0
1853	39216	42.9	0.0	7.0
2049	41657	42.9	0.0	7.0
5148	40188	42.9	0.0	7.0
16	44200	50.0	0.0	8.0
3619	43404	50.0	0.0	8.0
600	42108	57.1	, 0.0	7.0
4684	40125	57.1	0.0	7.0
1591	44634	62.5	0.0	8.0
1518	46399	71.4	0.0	7.0
1978	38827	71.4	0.0	7.0
17	44202	75.0	0.0	8.0
1975	41244	85.7	0.0	7.0
118	43970	87.5	0.0	8.0
114	43972	100.0	0.0	8.0

Table 13

A HOTO TO			
CloneID	ES No	ClusterID	SequenceName
M00007960D:E09	ES 168	142842	1513.A14.gz43 300454
M00007963D:D03	ES 168	142614	1513.A15.gz43 300470
M00021925A:H07	ES 168	120049	1513.D19.gz43 300537
M00022072A:E12	ES 168	153316	1513.E14.gz43_300458
M00022135D:D06	ES 168	145815	1513.F09.gz43_300379
M00022255B:F12	ES 168	158321	1513.H09.gz43_300381
M00022445D:E12	ES 168	101499	1513.J13.gz43_300447
M00022537D:C05	ES 168	168195	1513.L10.gz43 300401
M00022622A:G01	ES 168	99011	1513.N16.gz43_300499
M00022648C:D08	ES 168	169458	1513.O03.gz43_300292
M00022710C:H03	ES 168	171073	1513.P18.gz43_300533
M00023406A:G03	ES 168	189993	1521.C14.gz43_303619
M00027018B:F01	ES 168	94539	1521.F23.gz43_303766
M00027100A:D12	ES 168	220463	1521.H05.gz43_303480
M00027103D:B05	ES 168	189073	1521.H06.gz43_303496
M00027123D:F02	ES 168	186594	1521.H13.gz43_303608
M00027126C:H05	ES 168	222818	1521.H14.gz43_303624
M00027219B:G12	ES 168	188309	1521.J06.gz43_303498
M00027287B:H10	ES 168	217042	1521.K08.gz43_303531
M00027514C:F01	ES 168	187525	1521.N10.gz43_303566
M00027517C:F08	ES 168	215366	1521.N12.gz43_303598
M00027586B:B03	ES 168	218904	1521.O13.gz43_303615
M00027587C:F02	ES 168	185056	1521.O16.gz43_303663
M00027694C:C11	ES 168	186404	1521.P20.gz43_303728
M00028061D:D10	ES 168	432159	2128.A24.gz43_277785
M00028359D:F09	ES 168	188377	2128.H15.gz43_277648
M00028627B:F12	ES 168	427799	2128.K06.gz43_277507
M00032476D:F07	ES 168	155615	2128.P17.gz43_277688
M00032477A:B02	ES 168	125604	2128.P18.gz43_277704
M00032685B:C10	ES 168	427571	2130.H13.gz43_278017
M00032686C:D10	ES 168	38494	2130.H16.gz43_278065
M00032686D:G09	ES 168	78607	2130.H18.gz43_278097
M00032695B:A01	ES 168	44615	2130.I20.gz43_278130
M00032703D:E10	ES 168	90192	2130. <b>J</b> 18.gz43_278099
M00032732A:A03	ES 168	376753	2130.M21.gz43_278150
M00032736A:B06	ES 168	429735	2130.N02.gz43_277847
M00032738D:G11	ES 168	436888	2130.N09.gz43_277959
M00032745C:F03	ES 168	72838	2130.O05.gz43_277896
M00032766A:A10	ES 168	427907	2131.A01.gz43_307885
M00032779A:A04	ES 168	37875	2131.A19.gz43_308173
M00032780A:B09	ES 168	48238	2131,A23.gz43_308237
M00032783A:H08	ES 168	226324	2131.B04.gz43_307934
M00032786A:H04	ES 168	221686	2131.B14.gz43_308094
M00032809B:E10	ES 168	441801	2131.E06.gz43_307969

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Table 13

Table 15			
CloneID	ES No	ClusterID	SequenceName
M00032811A:G10	ES 168	48238	2131.E12.gz43 308065
M00032857A:B02	ES 168	34071	2131.I13.gz43_308085
M00032858D:H11	ES 168	26926	2131.I19.gz43_308181
M00032859C;E04	ES 168	441874	2131.I21.gz43 308213
M00032865A:D11	ES 168	440284	2131.J16.gz43 308134
M00032872B;A02	ES 168	62016	2131.K11.gz43 308055
M00032886A:D04	ES 168	48238	2131.M02.gz43_307913
M00032914B:D09	ES 168	12481	2131.P18.gz43_308172
M00032915B:D01	ES 168	37805	2131.P19.gz43_308188
M00039047C:A05	ES 168	376044	2152.A18.gz43_278491
M00039052B:H03	ES 168	36453	2152.B14.gz43_278428
M00039069D:E12	ES 168	387530	2152.D12.gz43_278398
M00039109A:H09	ES 168	375655	2152.H10.gz43_278370
M00039121C:C06	ES 168	375415	2152.J18.gz43_278500
M00039183B:E03	ES 168	400617	2176.A09.gz43_281808
M00039184A:A08	ES 168	400047	2176.A13.gz43 281872
M00039184C:C05	ES 168	402411	2176.A16.gz43 281920
M00039214C:D12	ES 168	390563	2153.D08.gz43_278834
M00039222C:G06	ES 168	375706	2153.F03.gz43 278756
M00039240A:H08	ES 168	390017	2153.I08.gz43_278839
M00039241A:H11	ES 168	376384	2153.I11.gz43_278887
M00039244C:F10	ES 168	379046	2153.J06.gz43_278808
M00039251A:G12	ES 168	372952	2153.K14.gz43_278937
M00039274C:D12	ES 168	377596	2153.O08.gz43_278845
M00039330B:H09	ES 168	60037	2154.G05.gz43_279173
M00039336A:C07	ES 168	376044	2154.G21.gz43_279429
M00039346C:G08	ES 168	390968	2154.I18.gz43_279383
M00039363B:H10	ES 168	409330	2176.B02.gz43_281697
M00039366C:B10	ES 168	400171	2176.B12.gz43_281857
M00039366D:A02	ES 168	404736	2176.B13.gz43_281873
M00039369D:D11	ES 168	407195	2176.C02.gz43_281698
M00039371A:E01	ES 168	407443	2176.C07.gz43 281778
M00039373B:B09	ES 168	403671	2176.C12.gz43_281858
M00039375A:C09	ES 168	411124	2176.C17.gz43_281938
M00039379A:D03	ES 168	411985	2176.D02.gz43_281699
M00039404B:G11	ES 168	377696	2154.M04.gz43 279163
M00039440C:F09	ES 168	376255	2155.A24.gz43_279855
M00039451B:C06	ES 168	389425	2155.B19.gz43_279776
M00039451B:F01	ES 168	378420	2155.B21.gz43_279808
M00039457A:D07	ES 168	373973	2155.C12.gz43_279665
M00039479A:A10	ES 168	376439	2155.F16.gz43_279732
M00039479D:C06	ES 168	375510	2155.F21.gz43_279812
M00039499C:G09	ES 168	413767	2176.E09.gz43_281812
M00039502D:H03	ES 168	401675	2176.E19.gz43_281972

Table 13

CloneID	ES No	ClusterID	SequenceName
M00039503B:G05	ES 168	413621	2176.F01.gz43 281685
M00039503D:E08	ES 168	412621	2176.F02.gz43_281701
M00039504B:F05	ES 168	402941	2176.F06.gz43 281765
M00039505B:G07	ES 168	413627	2176.F13.gz43_281877
M00039507D:E03	ES 168	412483	2176.F24.gz43 282053
M00039508D:C03	ES 168	411113	2176.G03.gz43 281718
M00039514A:G03	ES 168	402392	2176.G24.gz43 282054
M00039514B:E07	ES 168	407711	2176.H02.gz43 281703
M00039516C:H06	ES 168	400633	2176.H09.gz43 281815
M00039521A:D02	ES 168	400454	2176.H20.gz43 281991
M00039521C:B01	ES 168	405932	2176.H22.gz43 282023
M00039523A:D09	ES 168	402411	2176.I04.gz43 281736
M00039525B:H03	ES 168	409612	2176.I14.gz43 281896
M00039527B:D06	ES 168	406855	2176.I19.gz43 281976
M00039530B:H07	ES 168	414355	2176.J04.gz43_281737
M00039534A:H04	ES 168	49950	2176.J13.gz43_281881
M00039534D:E07	ES 168	412416	2176.J17.gz43 281945
M00039536B:E10	ES 168	402147	2176.J20.gz43 281993
M00039536B:H03	ES 168	413997	2176.J21.gz43_282009
M00039537B:F06	ES 168	401510	2176.J24.gz43_282057
M00039562A:D10	ES 168	400428	2176.K08.gz43_281802
M00039562D:B02	ES 168	400233	2176.K10.gz43_281834
M00039562D:G01	ES 168	408986	2176.K11.gz43_281850
M00039563C:D01	ES 168	407260	2176.K13.gz43_281882
M00039564B:F08	ES 168	408306	2176.K18.gz43_281962
M00039564C:H05	ES 168	409589	2176.K21.gz43_282010
M00039565B:H09	ES 168	402287	2176.K24.gz43_282058
M00039566D:E08	ES 168	400628	2176.L06.gz43_281771
M00039568C:E05	ES 168	94771	2176.L11.gz43_281851
M00039583A:G09	ES 168	408649	2176.L24.gz43_282059
M00039586A:C03	ES 168	406499	2176.M13.gz43_281884
M00039586D:D05	ES 168	406734	2176.M17.gz43_281948
M00039588A:H12	ES 168	202308	2176.M20.gz43_281996
M00039588B:H02	ES 168	409262	2176.M21.gz43_282012
M00039588D:F10	ES 168	402049	2176.M22.gz43_282028
M00039592C:F09	ES 168	413346	2176.N07.gz43_281789
M00039592D:E01	ES 168	401005	2176.N09.gz43_281821
M00039594A:F09	ES 168	413133	2176.N14.gz43_281901
M00039599A:E08	ES 168	407964	2176.O07.gz43_281790
M00039605D:E02	ES 168	401426	2176.P01.gz43_281695
M00039606C:B07	ES 168	202308	2176.P07.gz43_281791
M00039607A:F05	ES 168	402147	2176.P09.gz43_281823
M00039610A:G11	ES 168	413915	2176.P23.gz43_282047
M00039619C:B01	ES 168	376994	2155.I16.gz43_279735

Table 13

CloneID	ES No	ClusterID	SequenceName
M00039632A:C01	ES 168	60037	2155.J03.gz43 279528
M00039664B:H10	ES 168	376839	2155.N05.gz43_279564
M00039665C:B01	ES 168	.379805	2155.N15.gz43 279724
M00039674B:G11	ES 168	373680	2155.P11.gz43_279662
M00039675C;C05	ES 168	379879	2155.P22.gz43 279838
M00039679C:A02	ES 168	379279	2164.A24.gz43_280239
M00039681B:F05	ES 168	379046	2164.B15.gz43_280096
M00039698A:A06	ES 168	396785	2164.D18.gz43 280146
M00039730B:B07	ES 168	398028	2164.H18.gz43_280150
M00039734A:E06	ES 168	377772	2178.A12.gz43_282310
M00039752D:D07	ES 168	402070	2178.B16.gz43_282375
M00039753B:A11	ES 168	401748	2178.B17.gz43_282391
M00039760B:F12	ES 168	419751	2178.C15.gz43 282360
M00039760C:H07	ES 168	420504	2178.C18.gz43 282408
M00039774A:E11	ES 168	380025	2164.I10.gz43 280023
M00039778D:D05	ES 168	397167	2164.J03.gz43_279912
M00039820D:F08	ES 168	376074	2164.O24.gz43_280253
M00039823D:D09	ES 168	216179	2164.P10.gz43 280030
M00039871C:C01	ES 168	398061	2165.F09.gz43 280388
M00039883D:G06	ES 168	373905	2165.H01.gz43 280262
M00039885B:A10	ES 168	393635	2165.H06.gz43 280342
M00039893A:G12	ES 168	376808	2165.I10.gz43 280407
M00039905B:F09	ES 168	376773	2165.K01.gz43 280265
M00039977C:C05	ES 168	67549	2165.P01.gz43_280270
M00039978D:C04	ES 168	396969	2165.P07.gz43_280366
M00039986B:A11	ES 168	375655	2166.A06.gz43_281279
M00039988B:C08	ES 168	379879	2166.A11.gz43 281359
M00040001A:H02	ES 168	398831	2166.A23.gz43 281551
M00040070C:D11	ES 168	377696	2166.J11.gz43_281368
M00040076B:D01	ES 168	233814	2166.K07.gz43 281305
M00040096D:C03	ES 168	379879	2166.N07.gz43_281308
M00040113C:H09	ES 168	185432	2166.P24.gz43_281582
M00040134A:A07	ES 168	402411	2178.D07.gz43_282233
M00040141A:G10	ES 168	417259	2178.E04.gz43_282186
M00040145B:C12	ES 168	385980	2178.E21.gz43 282458
M00040160C:A04	ES 168	414821	2178.G05.gz43_282204
M00040161C:H06	ES 168	417426	2178.G11.gz43_282300
M00040161D:C03	ES 168	415527	2178.G12.gz43_282316
M00040171B:F01	ES 168	416819	2178.H19.gz43_282429
M00040171B:H03	ES 168	402353	2178.H20.gz43_282445
M00040181B:B06	ES 168	418340	2178.J12.gz43_282319
M00040181B:C05	ES 168	403837	2178.J13.gz43_282335
M00040183B:C06	ES 168	418682	2178.K02.gz43_282160
M00040183D:C08	ES 168	402534	2178.K05.gz43_282208

Table 13

A HIDIC TO			
CloneID	ES No	ClusterID	SequenceName
M00040188B:E05	ES 168	419255	2178.L07.gz43_282241
M00040189D:A06	ES 168	403154	2178.L12.gz43 282321
M00040190B:C02	ES 168	418482	2178.L14.gz43 282353
M00040196B:F10	ES 168	416914	2178.L20.gz43 282449
M00040197B:D05	ES 168	415950	2178.M03.gz43 282178
M00040222A:E06	ES 168	57183	2178.M22.gz43 282482
M00040222C:E06	ES 168	416370	2178.N03.gz43 282179
M00040223B:G05	ES 168	417240	2178.N07.gz43_282243
M00040227A:E07	ES 168	402070	2178.N16.gz43_282387
M00040230B:B01	ES 168	415058	2178.007.gz43_282244
M00040233C:F09	ES 168	416762	2178.O21.gz43_282468
M00040235C:D02	ES 168	402049	2178.P04.gz43_282197
M00039747A:H06	ES 169	423884	2184.A03.gz43_282699
M00039748A:E12	ES 169	422788	2184.A05.gz43_282731
M00039749B:G05	ES 169	402298	2184.A08.gz43_282779
M00039750C:F08	ES 169	403306	2184.A17.gz43_282923
M00039762C:D11	ES 169	140224	2184.A22.gz43_283003
M00039770C:B08	ES 169	424793	2184.B21.gz43_282988
M00039947A:G06	ES 169	423202	2184.C17.gz43_282925
M00039950C:C05	ES 169	403306	2184.D01.gz43_282670
M00039958D:D05	ES 169	402049	2184.D16.gz43_282910
M00040210D:C09	ES 169	424996	2184.F12.gz43_282848
M00040248A:G09	ES 169	402941	2178.P19.gz43_282437
M00040294B:E09	ES 169	402298	2184.G24.gz43_283041
M00040297B:G04	ES 169	423214	2184.H17.gz43_282930
M00040318C:B09	ES 169	403111	2184,J04.gz43_282724
M00040319A:E03	ES 169	422542	2184.J07.gz43_282772
M00040319C:F08	ES 169	235376	2184.J10.gz43_282820
M00040319D:G10	ES 169	423534	2184.J11.gz43_282836
M00040320A:B06	ES 169	401748	2184.J12.gz43_282852
M00040322B:A08	ES 169	420958	2184.K02.gz43_282693
M00040323D:F04	ES 169	423008	2184.K13.gz43_282869
M00040328D:A03	ES 169	402298	2184.L01.gz43_282678
M00040330D:C12	ES 169	421826	2184.L10.gz43_282822
M00040338C:D05	ES 169	396509	2184.M14.gz43_282887
M00040347B:B11	ES 169	424723	2184.N13.gz43_282872
M00040363A:G07	ES 169	403837	2184.O04.gz43_282729
M00040364B:E11	ES 169	422590	2184.O07.gz43_282777
M00042341A:D08	ES 169	0	1561.A04.gz43_314441
M00042341A:H04	ES 169	0	1561.A06.gz43_314473
M00042341D:G11	ES 169	0	1561.A14.gz43_314601
M00042342A:B04	ES 169	0	1561.A15.gz43_314617
M00042342C:H03	ES 169	0	1561.A18.gz43_314665

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00042342D:D03	ES 169	0	1561.A20.gz43_314697
M00042343A:C05	ES 169	0	1561.A24.gz43_314761
M00042343B:D05	ES 169	0	1561.B04.gz43_314442
M00042343B:E06	ES 169	0	1561.B05.gz43_314458
M00042343B:F12	ES 169	0	1561.B06.gz43_314474
M00042343D:F08	ES 169	0	1561.B09.gz43_314522
M00042344B:F02	ES 169	0	1561.B17.gz43_314650
M00042344B:H12	ES 169	0	1561.B19.gz43_314682
M00042344D:F03	ES 169	0	1561.C01.gz43_314395
M00042345A:E04	ES 169	0	1561.C03.gz43_314427
M00042345B:A05	ES 169	0	1561.C05.gz43_314459
M00042345D:B01	ES 169	0	1561.C10.gz43_314539
M00042345D:D04	ES 169	0	1561.C12.gz43_314571
M00042521A:E10	ES 169	0	1561.C16.gz43_314635
M00042521C:F11	ES 169	0	1561.C19.gz43_314683
M00042521D:G09	ES 169	0	1561.C22.gz43_314731
M00042522A:A05	ES 169	0	1561.C24.gz43_314763
M00042522C:H10	ES 169	0	1561.D10.gz43_314540
M00042522D:A08	ES 169	0	1561.D11.gz43_314556
M00042523B:H01	ES 169	0	1561.D20.gz43_314700
M00042523C:E08	ES 169	0	1561.D23.gz43_314748
M00042523C:H04	ES 169	0	1561.E02.gz43_314413
M00042523C:H06	ES 169	0	1561.E03.gz43_314429
M00042524C:C12	ES 169	0	1561.E15.gz43_314621
M00042524D:A10	ES 169	0	1561.E17.gz43_314653
M00042524D:D06	ES 169	0	1561.E19.gz43_314685
M00042525C:H07	ES 169	0	1561.F12.gz43_314574
M00042525D:E01	ES 169	0	1561.F15.gz43_314622
M00042526A:E10	ES 169	0	1561.F17.gz43_314654
M00042526A:F11	ES 169	0	1561,F19.gz43_314686
M00042526B:C12	ES 169	0	1561.F20.gz43_314702
M00042526C:B12	ES 169	0	1561.F23.gz43_314750
M00042526D:A02	ES 169	0	1561.F24.gz43_314766
M00042526D:A05	ES 169	0	1561.G01.gz43_314399
M00042526D:A07	ES 169	0	1561.G02.gz43_314415
M00042526D:D04	ES 169	0	1561.G04.gz43_314447
M00042527B:B01	ES 169	0	1561.G11.gz43_314559
M00042527C;A10	ES 169	0	1561.G14.gz43_314607
M00042527C:F01	ES 169	0	1561.G15.gz43_314623
M00042527D:E05	ES 169	0	1561,G20.gz43_314703
M00042527D:F12	ES 169	0	1561.G23.gz43_314751
M00042528C:G06	ES 169	0	1561,H07.gz43_314496
M00042528D:D09	ES 169	0	1561.H09.gz43_314528
M00042528D:H03	ES 169	0	1561.H10.gz43_314544

Table 13

Table 15	-		
CloneID	ES No	ClusterID	SequenceName
M00042529A:B12	ES 169	0	1561.H13.gz43_314592
M00042529A:G07	ES 169	0	1561.H15.gz43_314624
M00042529B:E03	ES 169	0	1561.H19.gz43_314688
M00042529D:D07	ES 169	0	1561.I02.gz43_314417
M00042530C:H02	ES 169	0	1561.I11.gz43_314561
M00042530D:A02	ES 169	0	1561.I12.gz43_314577
M00042530D:F09	ES 169	0	1561,I14.gz43_314609
M00042530D:H12	ES 169	0	1561.I17.gz43_314657
M00042531B:D12	ES 169	0	1561.I22.gz43_314737
M00042531B:G12	ES 169	0	1561.J01.gz43_314402
M00042531D:G08	ES 169	0	1561.J05.gz43_314466
M00042532A:D08	ES 169	0	1561.J09.gz43_314530
M00042532A:H03	ES 169	0	1561.J14.gz43_314610
M00042533B:F11	ES 169	0	1561.K02.gz43_314419
M00042533C:F04	ES 169	0	1561.K05.gz43_314467
M00042533D:B05	ES 169	0	1561.K06.gz43_314483
M00042534A:G10	ES 169	0	1561.K14.gz43_314611
M00042534B:B08	ES 169	0	1561.K16.gz43_314643
M00042534B:D06	ES 169	0	1561.K17.gz43_314659
M00042534C:A09	ES 169	0	1561.K19.gz43_314691
M00042534C:G04	ES 169	0	1561.K20.gz43_314707
M00042534D:D10	ES 169	0	1561.K23.gz43_314755
M00042535B:E04	ES 169	0	1561.L03.gz43_314436
M00042536B:G08	ES 169	0	1561.L15.gz43_314628
M00042536C:A06	ES 169	0	1561.L16.gz43_314644
M00042536C:B03	ES 169	0	1561.L18.gz43_314676
M00042537A:A07	ES 169	0	1561.L22.gz43_314740
M00042537A:D12	ES 169	0	1561.M01.gz43_314405
M00042537A:G09	ES 169	0	1561.M04.gz43_314453
M00042537A:H04	ES 169	0	1561.M05.gz43_314469
M00042537B:A07	ES 169	0	1561.M06.gz43_314485
M00042537B:B05	ES 169	0	1561.M07.gz43_314501
M00042537C:D04	ES 169	0	1561.M10.gz43_314549
M00042539B:D09	ES 169	0	1561.N06.gz43_314486
M00042539C:A04	ES 169	0	1561.N07.gz43_314502
M00042539C:E05	ES 169	0	1561.N08.gz43_314518
M00042539C:H08	ES 169	0	1561.N09.gz43_314534
M00042539D:A04	ES 169	0	1561.N10.gz43_314550
M00042540A:D11	ES 169	0	1561.N13.gz43_314598
M00042540A:G11	ES 169	0	1561,N14.gz43_314614
M00042540B:B03	ES 169	0	1561.N16.gz43_314646
M00042540C:A12	ES 169	0	1561.N17.gz43_314662
M00042540D:F12	ES 169	0	1561.N23.gz43_314758
M00042540D:H02	ES 169	0	1561.N24.gz43_314774

Table 13

CloneID	ES No	ClusterID	SequenceName
M00042541A;B07	ES 169	0	1561.O04.gz43_314455
M00042541A:E06	ES 169	0	1561.O06.gz43_314487
M00042541A:G02	ES 169	0	1561.O07.gz43 314503
M00042541B:B05	ES 169	0	1561.O09.gz43 314535
M00042542A:C11	ES 169	0	1561.O18.gz43 314679
M00042542B:B01	ES 169	0	1561.O21.gz43 314727
M00042542B:C11	ES 169	0	1561.O23.gz43_314759
M00042542B:E04	ES 169	0	1561.O24.gz43_314775
M00042543B:H12	ES 169	0	1561.P18.gz43_314680
M00042543C:F06	ES 169	0	1561.P21.gz43_314728
M00042546A:D03	ES 169	446572	1562.B01.gz43_207804
M00042546D:E06	ES 169	446389	1562.B13.gz43_207996
M00042547B:D11	ES 169	446531	1562.B22.gz43_208140
M00042548B;A01	ES 169	456845	1562.C12.gz43_207981
M00042548B:G01	ES 169	462149	1562.C14.gz43_208013
M00042550A:D12	ES 169	446674	1562.D12.gz43_207982
M00042550C:H10	ES 169	446981	1562.D18.gz43_208078
M00042552C:D02	ES 169	466280	1562.E22.gz43_208143
M00042552D:A11	ES 169	464091	1562.F01.gz43_207808
M00042554A:C02	ES 169	451780	1562.F04.gz43_207856
M00042554C:E02	ES 169	467262	1562.F08.gz43_207920
M00042554C:F09	ES 169	447597	1562.F10.gz43_207952
M00042554D:C08	ES 169	465594	1562.F12.gz43_207984
M00042555A:A04	ES 169	464067	1562.F17.gz43_208064
M00042555A:A10	ES 169	447152	1562.F18.gz43_208080
M00042555B;B07	ES 169	464905	1562.F20.gz43_208112
M00042555D:G10	ES 169	469511	1562.G05.gz43_207873
M00042556A:G12	ES 169	451737	1562.G10.gz43_207953
M00042556B:D12	ES 169	466719	1562.G12.gz43_207985
M00042556B;E10	ES 169	447174	1562.G14.gz43_208017
M00042557A:D09	ES 169	455075	1562.H05.gz43_207874
M00042557D:H12	ES 169	469837	1562.H17.gz43_208066
M00042558A:D03	ES 169	447517	1562.H18.gz43_208082
M00042558A:F11	ES 169	468330	1562.H20.gz43_208114
M00042558D:B07	ES 169	447308	1562.I03.gz43_207843
M00042560B:A01	ES 169	408386	1562.J01.gz43_207812
M00042560B:G10	ES 169	456462	1562.J04.gz43_207860
M00042561B:E12	ES 169	446620	1562.J21.gz43_208132
M00042561C:E12	ES 169	446621	1562.J24.gz43_208180
M00042562C:A07	ES 169	457405	1562.K14.gz43_208021
M00042563A:F10	ES 169	446829	1562.L02.gz43_207830
M00042563C:D08	ES 169	459536	1562.L09.gz43_207942
M00042563C:E02	ES 169	446614	1562.L10.gz43_207958
M00042563D:D02	ES 169	459523	1562.L11.gz43_207974

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CloneID	ES No	ClusterID	SequenceName
M00042563D:D09	ES 169	446495	1562.L12.gz43_207990
M00042564A:F01	ES 169	461559	1562.L19.gz43_208102
M00042564B:D11	ES 169	446495	1562.L22.gz43_208150
M00042564D:F10	ES 169	456471	1562.M06.gz43 207895
M00042565A:G05	ES 169	432159	1562,M10.gz43 207959
M00042565A:H03	ES 169	463487	1562.M12.gz43_207991
M00042567C:E02	ES 169	446703	1562.N16.gz43 208056
M00042567C:E07	ES 169	460972	1562.N18.gz43_208088
M00042567D.B08	ES 169	457783	1562.N21.gz43_208136
M00042567D:E12	ES 169	460493	1562.N24.gz43_208184
M00042568C:E03	ES 169	446657	1562.O14.gz43 208025
M00042568C:E08	ES 169	460766	1562.O15.gz43 208041
M00042569B:G07	ES 169	43338	1562.P03.gz43 207850
M00042569C:B05	ES 169	447346	1562.P05.gz43 207882
M00042569D:D02	ES 169	460190	1562.P09.gz43 207946
M00042570A:E08	ES 169	460516	1562.P11.gz43 207978
M00042570B:F11	ES 169	417078	1562.P18.gz43 208090
M00042570C:B12	ES 169	457842	1562.P21.gz43 208138
M00042571C:F03	ES 169	468783	1563.A14.gz43 208395
M00042571D:D06	ES 169	467057	1563,A15,gz43 208411
M00042572B:C07	ES 169	446302	1563.A19.gz43 208475
M00042572B:E05	ES 169	460789	1563.A20.gz43 208491
M00042573A:D05	ES 169	466971	1563,B05,gz43 208252
M00042573C:D05	ES 169	456052	1563,B14,gz43 208396
M00042573D:A10	ES 169	463896	1563.B21.gz43 208508
M00042574A:F05	ES 169	446409	1563.B23.gz43 208540
M00042574B:A07	ES 169	456925	1563.B24.gz43 208556
M00042574C:A04	ES 169	456920	1563,C04,gz43_208237
M00042575A:E01	ES 170	467327	1563.C14.gz43_208397
M00042575A:E02	ES 170	451383	1563.C15.gz43_208413
M00042575B:F02	ES 170	451382	1563.C20.gz43_208493
M00042575D:A11	ES 170	464275	1563.D01.gz43_208190
M00042575D:C05	ES 170	447417	1563.D03.gz43_208222
M00042576B:D11	ES 170	459961	1563.D12.gz43 208366
M00042576D:A08	ES 170	446213	1563.D17.gz43 208446
M00042576D:F01	ES 170	446839	1563.D21.gz43_208510
M00042577A:A11	ES 170	464547	1563.D23.gz43_208542
M00042577B;D05	ES 170	467051	1563.E06.gz43 208271
M00042578A:E08	ES 170	447597	1563.E14.gz43 208399
M00042579A:B05	ES 170	446933	1563.E22.gz43 208527
M00042579B:E05	ES 170	467644	1563.F01.gz43 208192
M00042580C:A03	ES 170	463951	1563.F13.gz43 208384
M00042691A:D08	ES 170	460244	1563.F19.gz43_208480

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00042691A:G04	ES 170	462767	1563.F21.gz43_208512
M00042691C:A05	ES 170	456985	1563,G04.gz43_208241
M00042691D:B03	ES 170	457890	1563.G08.gz43_208305
M00042691D:F03	ES 170	446776	1563,G12,gz43_208369
M00042692A:E11	ES 170	467293	1563.G16.gz43_208433
M00042692B:F04	ES 170	44503	1563.G22.gz43_208529
M00042693B:D03	ES 170	446549	1563.H09.gz43_208322
M00042694B:G05	ES 170	446962	1563,H18.gz43_208466
M00042694C:F01	ES 170	462008	1563.H23.gz43_208546
M00042695C:F05	ES 170	461734	1563.I09.gz43_208323
M00042695D:E11	ES 170	460884	1563.I12.gz43_208371
M00042695D:H01	ES 170	463368	1563.I14.gz43_208403
M00042697B:F06	ES 170	451454	1563.J01.gz43_208196
M00042697D:A03	ES 170	447190	1563.J03.gz43_208228
M00042698A:A01	ES 170	142559	1563.J05.gz43_208260
M00042699C:H06	ES 170	453605	1563.K08.gz43_208309
M00042700B:C12	ES 170	138049	1563.K12.gz43_208373
M00042700C:F11	ES 170	468109	1563.K20.gz43_208501
M00042700D:G08	ES 170	468979	1563.K24.gz43_208565
M00042702B:B07	ES 170	447326	1563.L20.gz43 208502
M00042702B:G07	ES 170	447826	1563.L22.gz43 208534
M00042702D:H01	ES 170	469944	1563.M02.gz43 208215
M00042704A:C02	ES 170	447433	1563.M18.gz43_208471
M00042705A:B07	ES 170	465339	1563.N08.gz43 208312
M00042706C:D02	ES 170	51939	1563.O02.gz43_208217
M00042707B:G05	ES 170	450929	1563.O14.gz43_208409
M00042707C:A09	ES 170	446922	1563.O16.gz43 208441
M00042709B:G05	ES 170	468930	1564.A01.gz43 296552
M00042709C:B05	ES 170	464937	1564.A02.gz43 296568
M00042710A:G10	ES 170	462393	1564.A08.gz43_296664
M00042710B:B09	ES 170	457975	1564.A09.gz43_296680
M00042711A;F01	ES 170	389425	1564,A20.gz43 296856
M00042711B;H04	ES 170	463143	1564.B01.gz43 296553
M00042711C:H10	ES 170	447006	1564.B06.gz43 296633
M00042711D;A03	ES 170	446191	1564.B07.gz43_296649
M00042711D;G04	ES 170	462604	1564.B09.gz43 296681
M00042712A:E08	ES 170	467780	1564.B12.gz43 296729
M00042712B:C04	ES 170	447429	1564.B16.gz43 296793
M00042712C:E06	ES 170	447649	1564.B19.gz43 296841
M00042713B:C08	ES 170	465785	1564.C01.gz43 296554
M00042713B:F03	ES 170	447189	1564.C03.gz43 296586
M00042713C:A09	ES 170	446230	1564.C04.gz43 296602
M00042713C:B08	ES 170	447268	1564.C05.gz43 296618
M00042713D:C06	ES 170	465782	1564.C07.gz43_296650

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Table 13

1 marc 10			
CloneID	ES No	ClusterID	SequenceName
M00042714A:G04	ES 170	456530	1564.C11.gz43_296714
M00042714A:H05	ES 170	447561	1564.C12.gz43_296730
M00042714B:C07	ES 170	447430	1564.C13.gz43_296746
M00042714B:E01	ES 170	447826	1564.C15.gz43_296778
M00042714C:C10	ES 170	465528	1564.C19.gz43_296842
M00042714C:F12	ES 170	449171	1564.C21.gz43_296874
M00042714D:E11	ES 170	452504	1564.C24.gz43_296922
M00042715D:A10	ES 170	457644	1564.D13.gz43_296747
M00042715D:E02	ES 170	446741	1564.D15.gz43_296779
M00042716A:F09	ES 170	446866	1564.D22.gz43_296891
M00042716D:G04	ES 170	462337	1564.E11.gz43_296716
M00042717B:G11	ES 170	462865	1564.E19.gz43_296844
M00042719A:H06	ES 170	452687	1564.F15.gz43_296781
M00042719C:H05	ES 170	405932	1564.F19.gz43_296845
M00042720C:E03	ES 170	461135	1564.G07.gz43_296654
M00042721A:G07	ES 170	446964	1564.G13.gz43_296750
M00042721B:A04	ES 170	446230	1564.G14.gz43_296766
M00042721D:B03	ES 170	457922	1564.G19.gz43_296846
M00042721D:D01	ES 170	450723	1564.G21.gz43_296878
M00042722A:G08	ES 170	462348	1564.H03.gz43_296591
M00042722B:D06	ES 170	460051	1564.H05.gz43_296623
M00042722D:C12	ES 170	459158	1564.H13.gz43_296751
M00042723A:D09	ES 170	446601	1564.H15.gz43_296783
M00042724A:G02	ES 170	453766	1564,I01.gz43_296560
M00042724D:B04	ES 170	446345	1564.I09.gz43_296688
M00042724D:H04	ES 170	447048	1564.I13.gz43_296752
M00042726B:E01	ES 170	446732	1564.I20.gz43_296864
M00042726D:G11	ES 170	462398	1564.J01.gz43_296561
M00042729A:F11	ES 170	461316	1564.K05.gz43_296626
M00042729A:H08	ES 170	462986	1564.K08.gz43_296674
M00042729B:F10	ES 170	461313	1564.K12.gz43_296738
M00042730D:D01	ES 170	424996	1564.L05.gz43_296627
M00042732A:A11	ES 170	453679	1564.L12.gz43_296739
M00042733C:C05	ES 170	454825	1564.M08.gz43_296676
M00042733D:G08	ES 170	450723	1564.M13.gz43_296756
M00042735A:G07	ES 170	469754	1564.M20.gz43_296868
M00042735A:G12	ES 170	469766	1564.M21.gz43_296884
M00042735B:A06	ES 170	403949	1564.M23.gz43_296916
M00042735C:G02	ES 170	420686	1564.N05.gz43_296629
M00042735D:A07	ES 170	463824	1564.N06.gz43_296645
M00042736A:F03	ES 170	447387	1564.N11.gz43_296725
M00042736B:G09	ES 170	447813	1564.N14.gz43_296773
M00042737A:A07	ES 170	463821	1564.N19.gz43_296853
M00042737B:C07	ES 170	390563	1564.O01.gz43_296566

Table 13

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CloneID	ES No	ClusterID	SequenceName
M00042737B;E02	ES 170	467189	1564.O02.gz43_296582
M00042737C:C07	ES 170	446740	1564.O04.gz43_296614
M00042737C:E03	ES 170	447579	1564.O05.gz43_296630
M00042737D:D03	ES 170	466614	1564.O08.gz43_296678
M00042738D:H12	ES 170	463217	1564.O19.gz43_296854
M00042739A;C06	ES 170	446242	1564.O20.gz43_296870
M00042739A:D02	ES 170	460078	1564.O21.gz43_296886
M00042739A:G07	ES 170	446922	1564.O23.gz43_296918
M00042739B:F06	ES 170	453762	1564.P02.gz43_296583
M00042740A:A01	ES 170	457656	1564.P09.gz43_296695
M00042742B:H03	ES 170	450940	1573.A06.gz43_208651
M00042744A:D11	ES 170	466894	1573.B07.gz43_208668
M00042745B;B06	ES 170	464622	1573.C01.gz43_208573
M00042745C:E11	ES 170	452399	1573.C05.gz43_208637
M00042745D:H04	ES 170	452093	1573.C08.gz43_208685
M00042747A:G12	ES 170	469150	1573.C22.gz43_208909
M00042747D:B01	ES 170	447293	1573.D01.gz43_208574
M00042747D:C08	ES 170	447421	1573.D02.gz43_208590
M00042748D:D08	ES 170	466920	1573.D10.gz43_208718
M00042750D:E07	ES 170	447645	1573.E04.gz43_208623
M00042881C:C11	ES 170	639372	1573.E15.gz43_208799
M00042882C:F06	ES 170	642146	1573.F10.gz43_208720
M00042882C:G07	ES 170	467293	1573.F12.gz43_208752
M00042882D:C04	ES 170	645690	1573.F15.gz43_208800
M00042883A:F06	ES 170	639849	1573.F18.gz43_208848
M00042884D:E03	ES 170	648467	1573.G17.gz43_208833
M00042885A:G09	ES 170	463060	1573.G21.gz43_208897
M00042886A:H03	ES 170	650364	1573.H04.gz43_208626
M00042886C:F01	ES 170	650195	1573.H11.gz43_208738
M00042886D:E10	ES 170	645470	1573.H16.gz43_208818
M00042887D;A11	ES 170	455996	1573.H21.gz43_208898
M00042888A:F02	ES 170	650231	1573.I03.gz43_208611
M00042889D:A01	ES 170	447161	1573.J08.gz43_208692
M00042889D:A12	ES 170	464205	1573.J10.gz43_208724
M00042890C:G11	ES 170	497434	1573.K01.gz43_208581
M00042890D:D03	ES 170	556711	1573.K03.gz43_208613
M00042890D:G05	ES 170	401426	1573.K06.gz43_208661
M00042891C:H01	ES 170	486238	1573.K19.gz43_208869
M00042892C:E03	ES 170	557974	1573.L09.gz43_208710
M00042892D:C04	ES 170	641890	1573.L10.gz43_208726
M00042892D:E06	ES 170	650852	1573.L13.gz43_208774
M00042892D:H04	ES 170	452506	1573.L14.gz43_208790
M00042894B:E05	ES 170	633946	1573.M08.gz43_208695
M00042894D:G05	ES 170	651029	1573.M17.gz43_208839

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CloneID	ES No	ClusterID	SequenceName
M00042895C:C10	ES 170	453908	1573,N01.gz43 208584
M00042896A:D04	ES 170	650756	1573.N12.gz43 208760
M00042896A;E03	ES 170	452611	1573.N13.gz43_208776
M00042896A:F09	ES 170	650944	1573.N14.gz43 208792
M00042898A:H05	ES 170	447141	1573.P02.gz43 208602
M00042900B:B10	ES 170	465104	1574.A03.gz43 208987
M00042900C:H11	ES 170	470593	1574.A04.gz43 209003
M00042901A:F12	ES 170	447750	1574.A11.gz43 209115
M00042901A:H11	ES 170	389591	1574.A14.gz43 209163
M00042902B:G02	ES 170	469608	1574.B04.gz43 209004
M00042902B:H01	ES 170	470462	1574.B05.gz43 209020
M00042902C:E11	ES 170	447660	1574.B07.gz43_209052
M00042903A:E04	ES 170	467710	1574.B12.gz43 209132
M00042903B:C09	ES 170	465984	1574.B15.gz43_209180
M00042904A:B01	ES 170	447212	1574.B24.gz43_209324
M00042904A:H10	ES 170	470641	1574.C03.gz43_208989
M00042905A:A07	ES 170	447147	1574.C07.gz43_209053
M00042905B:G03	ES 170	60260	1574.C11.gz43_209117
M00042905C:G08	ES 170	447815	1574.C14.gz43_209165
M00042906A:A12	ES 170	447218	1574.C17.gz43_209213
M00042906C:A10	ES 170	451624	1574.D04.gz43_209006
M00042907A:B11	ES 170	95617	1574.D14.gz43_209166
M00042907A:F03	ES 170	447692	1574.D15.gz43_209182
M00042907D:A11	ES 170	447173	1574.E02.gz43_208975
M00042908A:B01	ES 170	642691	1574.E06.gz43_209039
M00042908B:A11	ES 170	641069	1574.E12.gz43_209135
M00042908C:A03	ES 170	649744	1574.E18.gz43_209231
M00042908C:D12	ES 170	714629	1574.E20.gz43_209263
M00042908D:G12	ES 170	647086	1574.E24.gz43_209327
M00042909B:C04	ES 170	647639	1574.F07.gz43_209056
M00042909B:H08	ES 170	650398	1574.F12.gz43_209136
M00042909C:F10	ES 170	650235	1574.F17.gz43_209216
M00042909D:B11	ES 170	650564	1574.F18.gz43_209232
M00042910C:D03	ES 170	480508	1574.G04.gz43_209009
M00042910D:A02	ES 170	466697	1574.G06.gz43_209041
M00042910D:E11	ES 170	649965	1574.G08.gz43_209073
M00042911A:A02	ES 170	649810	1574.G11.gz43_209121
M00042911A:B02	ES 170	649900	1574.G12.gz43_209137
M00042911A:D04	ES 170	641029	1574.G15.gz43_209185
M00042911A:H12	ES 170	· 527355	1574.G18.gz43_209233
M00042911B:F10	ES 170	648855	1574.G21.gz43_209281
M00042911B:H08	ES 170	644376	1574.G23.gz43_209313
M00042911C:D01	ES 170	562247	1574.G24.gz43_209329
M00042912A:C01	ES 170	649965	1574.H06.gz43_209042

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CloneID	ES No	ClusterID	SequenceName
			33,4310
M00042912B:F11	ES 171	645924	1574.H11.gz43_209122
M00042912D:H08	ES 171	645954	1574.H16.gz43 209202
M00042912D:H10	ES 171	640356	1574.H17.gz43 209218
M00042913A:D09	ES 171	452989	1574.H18.gz43 209234
M00042913B:E10	ES 171	639740	1574.H22.gz43_209298
M00042914A:B05	ES 171	639371	1574.I09.gz43_209091
M00042914A:H10	ES 171	650423	1574.I16.gz43 209203
M00042914B:H03	ES 171	467364	1574.I20.gz43 209267
M00042914D:B10	ES 171	649852	1574.I22.gz43_209299
M00042915A:E06	ES 171	482043	1574.J05.gz43_209028
M00042915A:G10	ES 171	651051	1574.J06.gz43_209044
M00042915C:E05	ES 171	639923	1574.J12.gz43_209140
M00042915D:A11	ES 171	650527	1574.J15.gz43_209188
M00042915D:D06	ES 171	642151	1574.J17.gz43_209220
M00042915D:E03	ES 171	643940	1574.J18.gz43_209236
M00054791A:G04	ES 171	646404	1574.J21.gz43_209284
M00054791B:C09	ES 171	446974	1574.J24.gz43_209332
M00054792B:A03	ES 171	648159	1574.K18.gz43_209237
M00054793C:C01	ES 171	538237	1574.L07.gz43_209062
M00054793C:D11	ES 171	413767	1574.L11.gz43_209126
M00054793D:H11	ES 171	640306	1574.L13.gz43_209158
M00054794B:F09	ES 171	284269	1574.L23.gz43_209318
M00054794C:G11	ES 171	450506	1574.M06.gz43_209047
M00054794D:D02	ES 171	474298	1574.M09.gz43_209095
M00054794D:D08	ES 171	646568	1574.M10.gz43_209111
M00054795A:A08	ES 171	640181	1574.M16.gz43_209207
M00054796B:A01	ES 171	639391	1574.N04.gz43_209016
M00054796B:C08	ES 171	641525	· 1574.N05.gz43_209032
M00054797C:F03	ES 171	644789	1574.N20.gz43_209272
M00054797D:F01	ES 171	472101	1574.O01.gz43_208969
M00054798B:A01	ES 171	453572	1574.006.gz43_209049
M00054798D:A12	ES 171	641315	1574.O15.gz43_209193
M00054798D:F01	ES 171	650773	1574.O17.gz43_209225
M00054799C:G11	ES 171	641875	1574.P02.gz43_208986
M00054799D:H02	ES 171	645223	1574.P05.gz43_209034
M00054800B:C06	ES 171	639787	1574.P10.gz43_209114
M00054800B:C11	ES 171	513262	1574.P12.gz43_209146
M00054800B:E08	ES 171	474298	1574.P15.gz43_209194
M00054800D:D08	ES 171	639901	1574.P23.gz43_209322
M00054800D:F08	ES 171	513248	1575.A01.gz43_209339
M00054911A:C08	ES 171	549699	1575,A05.gz43_209403
M00054911A:G01	ES 171	559776	1575,A10.gz43_209483
M00054911A:H06	ES 171	528616	1575.A13.gz43_209531

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Table 15			
CloneID	ES No	ClusterID	SequenceName
M00054911B:E10	ES 171	639992	1575.A15.gz43_209563
M00054911D:D03	ES 171	41878	1575.B02.gz43_209356
M00054912A:H08	ES 171	646949	1575.B05.gz43_209404
M00054912B:C05	ES 171	589098	1575.B08.gz43_209452
M00054912C:C01	ES 171	641057	1575.B13.gz43_209532
M00054912D:F11	ES 171	182437	1575.B19.gz43_209628
M00054912D:G01	ES 171	417130	1575.B20.gz43_209644
M00054912D:G04	ES 171	640230	1575.B21.gz43_209660
M00054913A:B12	ES 171	377855	1575.B24.gz43_209708
M00054913C:G03	ES 171	451811	1575.C12.gz43_209517
M00054914C:D12	ES 171	447246	1575.D04.gz43_209390
M00054914D:G07	ES 171	467381	1575.D09.gz43_209470
M00054915A:G03	ES 171	413767	1575.D13.gz43_209534
M00054915D:B08	ES 171	452220	1575.D21.gz43_209662
M00054915D:D05	ES 171	646293	1575.D22.gz43_209678
M00054915D:E07	ES 171	468109	1575.D24.gz43 209710
M00054916A:A05	ES 171	649872	1575.E02.gz43 209359
M00054916A;E05	ES 171	639256	1575.E05.gz43 209407
M00054916A:F10	ES 171	639394	1575.E06.gz43 209423
M00054916B:E02	ES 171	648664	1575.E11.gz43 209503
M00054916C:C04	ES 171	545980	1575.E13.gz43_209535
M00054917A:F07	ES 171	473854	1575,E22.gz43 209679
M00054917B:A05	ES 171	648609	1575.E23.gz43 209695
M00054917B:F01	ES 171	649082	1575.F03.gz43_209376
M00054917C:D03	ES 171	643843	1575.F08.gz43 209456
M00054917C:F03	ES 171	648532	1575.F10.gz43 209488
M00054917D:A03	ES 171	647991	1575.F14.gz43 209552
M00054917D:A12	ES 171	644692	1575.F16.gz43 209584
M00054917D:D12	ES 171	639662	1575.F18.gz43 209616
M00054917D:E05	ES 171	639255	1575.F19.gz43 209632
M00054917D:H02	ES 171	648532	1575.F23.gz43_209696
M00054918A:D02	ES 171	649506	1575.G02.gz43 209361
M00054918A:F09	ES 171	553100	1575.G04.gz43 209393
M00054918B:H01	ES 171	449861	1575.G10.gz43 209489
M00054918B:H08	ES 171	645252	1575,G11.gz43 209505
M00054918D:C03	ES 171	456923	1575.G15.gz43 209569
M00054918D:C11	ES 171	452204	1575.G16.gz43_209585
M00054918D:G02	ES 171	645252	1575,G18.gz43 209617
M00054918D:H09	ES 171	468222	1575,G19.gz43 209633
M00054919A:H04	ES 171	452845	1575.G24.gz43 209713
M00054919C:F06	ES 171	644556	1575.H05.gz43_209410
M00054919D:H12	ES 171	642084	1575,H14.gz43 209554
M00054919B:H12	ES 171	61616	1575.H16.gz43_209586
M00054920A:B07	ES 171	554010	1575.H18.gz43_209618
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Table 13

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M00054920A:C11	ES 171	564440	1575.H22.gz43_209682
M00054920B:C04	ES 171	649933	1575.I03.gz43_209379
M00054920B:C12	ES 171	500337	1575.I05.gz43_209411
M00054920C:A06	ES 171	643843	1575.I10.gz43_209491
M00054920C:D05	ES 171	452707	1575.I12.gz43_209523
M00054920C:F02	ES 171	649555	1575.I15.gz43_209571
M00055426B:B02	ES 171	452986	1575.J06.gz43_209428
M00055426C:B12	ES 171	447574	1575.J14.gz43_209556
M00055426C:C10	ES 171	649746	1575.J15.gz43_209572
M00055426D:F03	ES 171	639144	1575.J21.gz43_209668
M00055427A:F01	ES 171	643924	1575.K05.gz43_209413
M00055427A:F02	ES 171	654723	1575.K06.gz43_209429
M00055427B:E01	ES 171	41141	1575.K10.gz43_209493
M00055427B:F06	ES 171	640814	1575.K11.gz43_209509
M00055427C:A06	ES 171	504568	1575.K13.gz43_209541
M00055427C:E12	ES 171	630269	1575.K17.gz43_209605
M00055427C:F07	ES 171	650487	1575.K18.gz43_209621
M00055427C:H11	ES 171	468783	1575.K19.gz43_209637
M00055427D:E05	ES 171	456420	1575.K21.gz43_209669
M00055428A:C02	ES 171	643279	1575.L01.gz43_209350
M00055428B:H02	ES 171	645347	1575,L05.gz43_209414
M00055428C:G06	ES 171	243722	1575.L12.gz43_209526
M00055428D:G12	ES 171	611927	1575.L18.gz43_209622
M00055429A:H04	ES 171	639520	1575.L22.gz43_209686
M00055429B:B12	ES 171	467989	1575.M01.gz43_209351
M00055429B:E12	ES 171	446254	1575.M02.gz43_209367
M00055429B:G04	ES 171	639444	1575.M03.gz43_209383
M00055429B:H02	ES 171	447254	1575.M04.gz43_209399
M00055429D:G07	ES 171	584071	1575.M11.gz43_209511
M00055430B:E08	ES 171	495143	1575.M16.gz43_209591
M00055430B:H02	ES 171	560700	1575.M19.gz43_209639
M00055430C:G11	ES 171	452293	1575.M23.gz43_209703
M00055430D:F04	ES 171	650184	1575.N01.gz43_209352
M00055431A:E01	ES 171	451994	1575.N05.gz43 209416
M00055431A:H05	ES 171	645344	1575.N07.gz43_209448
M00055431B:A01	ES 171	570812	1575,N09.gz43 209480
M00055431C:H08	ES 171	639523	1575.N15.gz43_209576
M00055432A:A03	ES 171	446495	1575.N19.gz43 209640
M00055432A:D05	ES 171	639150	1575.N21.gz43_209672
M00055432B:B04	ES 171	642073	1575.N23.gz43_209704
M00055432B:H02	ES 171	641542	1575.002.gz43_209369
M00055432C:D12	ES 171	447913	1575.005.gz43_209417
M00055432C:F01	ES 171	645162	1575.007.gz43_209449
M00055432D:H12	ES 171	655551	1575.015.gz43_209577

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CloneID	ES No	ClusterID	SequenceName
M00055433A:B07	ES 171	644371	1575.O16.gz43_209593
M00055433A:B08	ES 171	446932	1575.O17.gz43_209609
M00055433A:C02	ES 171	641925	1575.O18.gz43_209625
M00055433A:E04	ES 171	649195	1575.O22.gz43_209689
M00055433C:A11	ES 171	638758	1575.O24.gz43_209721
M00055433D;A09	ES 171	651211	1575.P02.gz43_209370
M00055433D:C07	ES 171	458734	1575.P04.gz43_209402
M00055433D:F06	ES 171	492242	1575.P06.gz43_209434
M00055434A:A03	ES 171	647109	1575.P08.gz43_209466
M00055434C:B11	ES 171	651020	1575.P18.gz43_209626
M00055434D:B06	ES 171	640369	1575.P21.gz43_209674
M00055434D;E09	ES 171	57183	1575.P22.gz43_209690
M00055435B:C09	ES 171	639829	1576.A04.gz43_209771
M00055435C:E12	ES 171	640055	1576.A08.gz43_209835
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M00055436B:B06	ES 171	639711	1576.A19.gz43_210011
M00055436B:B09	ES 171	454409	1576.A20.gz43_210027
M00055437D:B06	ES 171	466092	1576.B17.gz43_209980
M00055439B:B07	ES 171	640522	1576.C21.gz43_210045
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M00055439D:E08	ES 171	648506	1576.D11.gz43_209886
M00055440D:D02	ES 171	653616	1576.E09.gz43_209855
M00055441A:G07	ES 171	640734	1576.E16.gz43_209967
M00055441C:H06	ES 171	640230	1576.F03.gz43_209760
M00055443D:G04	ES 171	649288	1576.G13.gz43_209921
M00055444D:C05	ES 171	639750	1576.H03.gz43_209762
M00055446B:A12	ES 171	484145	1576.I05.gz43_209795
M00055446B:D08	ES 171	639928	1576.I08.gz43_209843
M00055450B:G07	ES 171	526606	1576.L08.gz43_209846
M00055451B:D08	ES 171	446439	1576.M09.gz43_209863
M00055451B:D12	ES 171	456125	1576.M10.gz43_209879
M00055451C:B08	ES 171	624440	1576.M12.gz43_209911
M00055451C:E10	ES 171	639786	1576.M14.gz43_209943
M00055453D:E12	ES 171	560791	1576.O12.gz43_209913
M00055455B:G12	ES 171	479051	1576.P24.gz43_210106
M00055456A:B03	ES 171	559369	1585.A06.gz43_210283
M00055456A:F01	ES 171	642478	1585.A10.gz43_210347
M00055456B:G08	ES 171	641645	1585.A22.gz43_210539
M00055456C:A10	ES 171	466440	1585.B02.gz43_210220
M00055456C:G04	ES 171	484126	1585.B06,gz43_210284
M00055456D:E06	ES 171	452202	1585.B13.gz43_210396
M00055456D:G04	ES 171	639932	1585.B14.gz43_210412
M00055457D:F09	ES 171	650605	1585,B23.gz43_210556

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M00055458D:F04	ES 171	640997	1585.C16.gz43 210445
M00055459B:A02	ES 171	641467	1585.C22.gz43 210541
M00055460A:D01	. ES 171	640662	1585.D14.gz43 210414
M00055460B:G06	ES 171	598589	1585.E03.gz43 210239
M00055460C:C12	ES 171	640561	1585.E06.gz43_210287
M00055460C:D10	ES 171	592346	1585.E07.gz43 210303
M00055460C:G09	ES 171	592346	1585.E11.gz43_210367
M00055460D:B06	ES 171	641066	1585.E15.gz43_210431
M00055461A:A06	ES 171	643991	1585.E23.gz43_210559
M00055461A.H03	ES 171	447863	1585.F03.gz43_210240
M00055461C:E05	ES 171	640792	1585.F13.gz43_210400
M00055461D:C09	ES 171	607422	1585.F17.gz43_210464
M00055462A;A09	ES 171	640368	1585.F22.gz43_210544
M00055462C:A11	ES 172	640400	1585.G12.gz43_210385
M00055462C:C03	ES 172	598589	1585.G14.gz43_210417
M00055462D:H12	ES 172	412416	1585.G22.gz43_210545
M00055463A:A11	ES 172	398061	1585.G23.gz43_210561
M00055463D:G01	ES 172	544461	1585.H09.gz43_210338
M00055464B:E06	ES 172	640913	1585.H13.gz43_210402
M00055464B:E11	ES 172	556654	1585.H14.gz43_210418
M00055464B:G03	ES 172	664711	1585.H15.gz43_210434
M00055464D:A04	ES 172	661194	1585.H19.gz43_210498
M00055464D:F08	ES 172	650914	1585.H22.gz43_210546
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M00055466A:C05	ES 172	549611	1585.I15.gz43_210435
M00055466C:A01	ES 172	559343	1585.I23.gz43_210563
M00055467A:A07	ES 172	555820	1585.J07.gz43_210308
M00055467D:A01	ES 172	559549	1585.J20.gz43_210516
M00055467D:C10	ES 172	51939	1585.J22.gz43_210548
M00055467D:G08	ES 172	505933	1585.J24.gz43_210580
M00055468A:A05	ES 172	646318	1585.K02.gz43_210229
M00055468C:B07	ES 172	639726	1585.K16.gz43_210453
M00055469A:D08	ES 172	640068	1585.K24.gz43_210581
M00055469B:A06	ES 172	553402	1585.L04.gz43_210262
M00055469C:F09	ES 172	645201	1585.L12.gz43_210390
M00055470C:F03	ES 172	562067	1585.M07.gz43_210311
M00055470D:F12	ES 172	640293	1585.M12.gz43_210391
M00055491A:H01	ES 172	584745	1585.M17.gz43_210471
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M00055492A:H06	ES 172	646303	1585.N10.gz43_210360

Table 13

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M00055493D:B07	ES 172	555193	1585.O14.gz43_210425
M00055493D:D12	ES 172	650353	1585.O16.gz43 210457
M00055494A;A06	ES 172	645117	1585.O18.gz43_210489
M00055494C:G10	ES 172	562236	1585.P05.gz43_210282
M00055494C:G11	ES 172	558839	1585.P06.gz43_210298
M00055495A:D11	ES 172	642415	1585.P12.gz43_210394
M00055495C:F03	ES 172	237288	1585.P22.gz43_210554
M00055516B:E11	ES 172	552783	1587.A04.gz43_211407
M00055517A:D09	ES 172	650161	1587,A16,gz43_211599
M00055517C:H07	ES 172	492483	1587.B02.gz43_211376
M00055517D:D09	ES 172	446984	1587.B06.gz43_211440
M00055517D:D11	ES 172	290226	1587.B07.gz43_211456
M00055519A:C01	ES 172	558785	1587.B23.gz43_211712
M00055519A:F08	ES 172	234606	1587.C01.gz43_211361
M00055519A:H01	ES 172	644051	1587.C03.gz43_211393
M00055520B:D11	ES 172	590218	1587.D04.gz43_211410
M00055520B:E04	ES 172	481220	1587.D05.gz43_211426
M00055521C:B08	ES 172	597780	1587.E01.gz43_211363
M00055521C:B09	ES 172	638943	1587.E02.gz43_211379
M00055521C:C08	ES 172	639056	1587.E03.gz43_211395
M00055521C:D02	ES 172	559324	1587.E04.gz43_211411
M00055522C:F06	ES 172	561202	1587.F04.gz43_211412
M00055523C:F05	ES 172	551518	1587.F21.gz43_211684
M00055524B:B08	ES 172	640419	1587.G08.gz43_211477
M00055524B:D11	ES 172	648747	1587.G10.gz43_211509
M00055528A:E08	ES 172	641615	1587.J11.gz43_211528
M00055528D:B02	ES 172	645544	1587.K05.gz43_211433
M00055528D:H07	ES 172	649932	1587.K13.gz43_211561
M00055529D:D05	ES 172	643723	1587.L03.gz43_211402
M00055529D:D11	ES 172	646711	1587.L04.gz43_211418
M00055529D:G03	ES 172	555326	1587.L06.gz43_211450
M00055530A:C07	ES 172	649149	1587.L09.gz43_211498
M00055531B:D10	ES 172	639132	1587,M04.gz43_211419
M00055531B:E05	ES 172	650617	1587.M05.gz43_211435
M00055531C:C04	ES 172	644479	1587.M10.gz43_211515
M00055531D:E06	ES 172	639213	1587.M13.gz43_211563
M00055532C:G08	ES 172	639459	1587.N03.gz43_211404
M00055532D:A12	ES 172	207552	1587.N04.gz43_211420
M00055533B:B11	ES 172	641216	1587.N15.gz43_211596
M00055533D:G02	ES 172	116869	1587.O04.gz43_211421
M00055534A:E06	ES 172	194095	1587.O05.gz43_211437

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Table 13			
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M00055536D:D12	ES 172	447869	1588.A07.gz43_217345
M00055537C:A01	ES 172	552001	1588.A11.gz43_211923
M00055537C:A01	ES 172	552001	1588.A11.gz43_217409
M00055537C:E04	ES 172	451671	1588.A15.gz43_217473
M00055538A:C05	ES 172	419706	1588.A18.gz43_212035
M00055538B:G09	ES 172	639461	1588.A21.gz43_217569
M00055538C:E04	ES 172	642631	1588.B02.gz43_211780
M00055538D:D12	ES 172	649335	1588.B06.gz43_211844
M00055538D:D12	ES 172	649335	1588.B06.gz43_217330
M00055538D:F12	ES 172	640089	1588.B09.gz43_211892
M00055538D:H07	ES 172	648206	1588.B11.gz43_211924
M00055540D:G11	ES 172	403419	1588.C19.gz43_217539
M00055541C:D02	ES 172	643277	1588.D05.gz43 211830
M00055542B:B11	ES 172	453606	1588.D17.gz43 212022
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M00055542C:B02	ES 172	451361	1588.D21.gz43 217572
M00055542C:D07	ES 172	639114	1588.D22.gz43 212102
M00055542C:H05	ES 172	641031	1588.E02.gz43 217269
M00055543B:A06	ES 172	568331	1588.E15.gz43_211991
M00055543C:B09	ES 172	638917	1588.E16.gz43 217493
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M00055543D:H03	ES 172	639543	1588,F02,gz43 211784
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M00055545A:C01	ES 172	644173	1588.F23.gz43 217606
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M00055547B:C05	ES 172	557983	1588.H21.gz43 212090
M00055547B:G09	ES 172	648494	1588.H24.gz43 217624
M00055547C:B01	ES 172	448673	1588,I02,gz43 211787
M00055547C:B07	ES 172	650982	1588.I04.gz43 217305
M00055547D:G06	ES 172	641401	1588.I15.gz43_217481

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M00055553C:D06
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M00055553D:C07         ES 172         640747         1588.N16.gz43_21201           M00055553D:E06         ES 172         645781         1588.N18.gz43_21735           M00055554C:B04         ES 172         645781         1588.005.gz43_21732           M00055554D:F01         ES 172         644012         1588.005.gz43_21732           M00055554D:F01         ES 172         377692         1588.012.gz43_21745           M00055555D:B05         ES 172         639395         1588.022.gz43_21756
M00055553D:E06         ES 172         645781         1588.N18.ga43_21753           M00055554C:B04         ES 172         644012         1588.005.gz43_21732           M000555554D:F01         ES 172         377692         1588.001.gz43_21742           M00055555D:B05         ES 172         639395         1588.022.gz43_21755
M00055554C:B04         ES 172         644012         1588.005.gz43_21732           M00055554D:F01         ES 172         377692         1588.011.gz43_21742           M00055555D:B05         ES 172         639395         1588.022.gz43_21759
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M00055556A:A04 ES 172 646350 1588 P05 gz43 21184
M00055556A:E07 ES 172 649099 1588.P06.gz43_21185
M00055556A:E07 ES 172 649099 1588.P06.gz43_21734
M00055556C:H09 ES 172 701221 1588.P14.gz43_21747
M00055557A:A04 ES 172 447023 1588.P18.gz43_21753
M00055557A:C11 ES 172 639425 1588.P21.gz43_21209
M00055557B:B10 ES 172 477064 1588 P24.gz43_21763
M00055557B:F07 ES 172 509798 1597.A03.gz43_21217
M00055558B:G11 ES 172 637966 1597.A16.gz43_21238
M00055558D:C08 ES 172 460666 1597.A21.gz43_21246
M00055558D:D07 ES 172 645913 1597.A23.gz43_21249
M00055559A:D06 ES 172 469731 1597.B06.gz43_21222

Table 13

Table 15			
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M00055561B:G12	ES 172	645538	1597.C24.gz43 212517
M00055561C:C07	ES 172	639781	1597.D05.gz43_212214
M00055562B:D03	ES 172	592122	1597.D14.gz43 212358
M00055562B:H02	ES 172	475152	1597.D17.gz43 212406
M00055562C:F02	ES 172	640195	1597.D22.gz43_212486
M00055562D:B06	ES 172	639056	1597.E05.gz43_212215
M00055563A:A02	ES 172	640356	1597.E08.gz43 212263
M00055563A:C03	ES 172	647431	1597.E09.gz43 212279
M00055563A:D02	ES 172	449258	1597.E10.gz43_212295
M00055565A:C08	ES 172	454825	1597.F10.gz43 212296
M00055565A:F11	ES 172	553318	1597.F16.gz43 212392
M00055565B:F04	ES 172	596882	1597.F18.gz43 212424
M00055565D:G08	ES 172	664195	1597.G03.gz43_212185
M00055566A:E02	ES 172	639593	1597.G06.gz43 212233
M00055567A:A06	ES 172	446230	1597.G23.gz43 212505
M00055567A:A11	ES 172	647069	1597.G24.gz43_212521
M00055567B:G07	ES 172	649829	1597.H04.gz43_212202
M00055568A:B04	ES 172	556511	1597.H15.gz43 212378
M00055568A:D05	ES 172	646293	1597.H17.gz43 212410
M00055568C:F07	ES 172	638869	1597.H24.gz43 212522
M00055568C:G11	ES 172	641890	1597.I01.gz43 212155
M00055569A:C08	ES 172	643488	1597.I10.gz43_212299
M00055569A:D01	ES 172	650740	1597.I12.gz43 212331
M00055569A:F06	ES 172	644928	1597.I18.gz43 212427
M00055569B:G10	ES 172	553114	1597.I24.gz43 212523
M00055569D:C02	ES 172	557975	1597.J06.gz43 212236
M00055570B:F07	ES 172	659031	1597.J12.gz43 212332
M00055572C:E08	ES 172	447423	1597.L05.gz43_212222
M00055573A:B11	ES 172	666644	1597.L12.gz43_212334
M00055573B:D11	ES 172	589483	1597.L18.gz43 212430
M00055573B:E01	ES 172	553602	1597.L19.gz43 212446
M00055574A:D12	ES 172	642527	1597.M09.gz43 212287
M00055574A:F12	ES 172	640181	1597.M11.gz43 212319
M00055574B:D04	ES 172	557713	1597.M15.gz43 212383
M00055574B:F06	ES 172	650235	1597.M17.gz43 212415
M00055574D:B07	ES 172	649431	1597.M24.gz43 212527
M00055574D:E02	ES 173	649668	1597.N02.gz43 212176
M00055575C:B04	ES 173	45921	1597.N14.gz43 212368
M00055575D:G04	ES 173	640956	1597.N18.gz43 212432
M00055576A:D09	ES 173	650593	1597.N21.gz43 212480

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Table 13

Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055576A:F07	ES 173	649349	1597.N23.gz43_212512
M00055576D:E09	ES 173	643162	1597.O12.gz43_212337
M00055577A:G09	ES 173	640851	1597.O20.gz43_212465
M00055577B:F05	ES 173	648483	1597.024.gz43_212529
M00055577C:G02	ES 173	639698	1597.P05.gz43_212226
M00055578B:D05	ES 173	362177	1597.P12.gz43_212338
M00055578C:F11	ES 173	668852	1597.P22.gz43_212498
M00055578D:E11	ES 173	561602	1597.P24.gz43_212530
M00055579C:D04	ES 173	641191	1598.A07.gz43_212627
M00055579D:C11	ES 173	459521	1598.A11.gz43_212691
M00055579D:G09	ES 173	641957	1598.A13.gz43_212723
M00055580B:B08	ES 173	641425	1598.A17.gz43_212787
M00055581A:C02	ES 173	640025	1598.B02.gz43_212548
M00055581C:A01	ES 173	648580	1598.B04.gz43_212580
M00055581C:B08	ES 173	647360	1598.B05.gz43_212596
M00055582B:A06	ES 173	648159	1598.B15.gz43_212756
M00055583A:A05	ES 173	641262	1598.C05.gz43_212597
M00055583C:A01	ES 173	86311	1598.C18.gz43_212805
M00055584A;G11	ES 173	641925	1598.D06.gz43_212614
M00055584B:B01	ES 173	639070	1598.D07.gz43 212630
M00055585A:E12	ES 173	65 1000	1598.D21.gz43 212854
M00055585B:F01	ES 173	415538	1598.E02.gz43 212551
M00055585C:F05	ES 173	449247	1598.E06.gz43 212615
M00055586A:F05	ES 173	639116	1598.E11.gz43 212695
M00055586C:A06	ES 173	553087	1598.E21.gz43 212855
M00055586D:G07	ES 173	645707	1598.F05.gz43 212600
M00055588A:C03	ES 173	639444	1598.G05.gz43 212601
M00055588B:H11	ES 173	140909	1598.G11.gz43 212697
M00055588C:G09	ES 173	509973	1598.G15.gz43 212761
M00055589A:B06	ES 173	646914	1598.G21.gz43 212857
M00055589B:E08	ES 173	644572	1598.H03.gz43 212570
M00055589B:H02	ES 173	642008	1598.H04.gz43 212586
M00055590A:B03	ES 173	464091	1598.H11.gz43 212698
M00055590A:E01	ES 173	645264	1598.H15.gz43_212762
M00055590D:G07	ES 173	635439	1598.I11.gz43 212699
M00055591A;B08	ES 173	641472	1598.I12.gz43 212715
M00055591D:A07	ES 173	556868	1598,J04.gz43 212588
M00055591D:E08	ES 173	512521	1598.J05.gz43 212604
M00055592B:C10	ES 173	641467	1598.J12.gz43 212716
M00055593A:F08	ES 173	641838	1598.K09.gz43 212669
M00055593C:D08	ES 173	638941	1598.K14.gz43 212749
M00055594B:A01	ES 173	446371	1598.K21.gz43 212861
M00055594C:B03	ES 173	470769	1598.L04.gz43 212590
M00055594C:F11	ES 173	646590	1598.L06.gz43 212622
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Table 13

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M00055595C:G03	ES 173	478229	1598.M02.gz43 212559
M00055595D:C04	ES 173	554273	1598.M04.gz43 212591
M00055596A:C02	ES 173	144626	1598.M08.gz43 212655
M00055596A:E07	ES 173	640826	1598.M10.gz43 212687
M00055596B:D03	ES 173	458979	1598.M16.gz43 212783
M00055597B:B04	ES 173	463028	1598,N05,gz43 212608
M00055597D:B05	ES 173	641440	1598.N09.gz43 212672
M00055597D:E10	ES 173	564440	1598.N11.gz43 212704
M00055599D:C08	ES 173	640298	1598.N23.gz43_212896
M00055600A;C04	ES 173	507188	1598.O07.gz43 212641
M00055600C:C02	ES 173	647312	1598.O17.gz43 212801
M00055600D:B02	ES 173	515931	1598.O19.gz43 212833
M00055601B:D12	ES 173	640695	1598,P05.gz43 212610
M00055601B:H02	ES 173	641191	1598,P07.gz43 212642
M00055601C:C11	ES 173	553925	1598.P10.gz43 212690
M00055602A:D04	ES 173	640634	1598.P20.gz43 212850
M00055602B:B10	ES 173	201904	1599.A01.gz43 212923
M00055602B:B12	ES 173	644764	1599.A03.gz43_212955
M00055602D:G08	ES 173	643906	1599.A15.gz43_213147
M00055603D:A09	ES 173	641338	1599,B01.gz43_212924
M00055604D:E07	ES 173	643279	1599.B09.gz43_213052
M00055604D:F05	ES 173	559380	1599,B10,gz43_213068
M00055606A:B11	ES 173	554833	1599.B19.gz43_213212
M00055606A;F09	ES 173	461524	1599.B21.gz43_213244
M00055606C:F04	ES 173	641839	1599,C09,gz43_213053
M00055606D:C05	ES 173	641542	1599.C12.gz43_213101
M00055608C:E03	ES 173	641680	1599.D10.gz43_213070
M00055608C:G11	ES 173	607715	1599.D15.gz43_213150
M00055609B:D10	ES 173	460929	1599.E01.gz43_212927
M00055609B:F10	ES 173	640930	1599.E03.gz43_212959
M00055610B:E04	ES 173	454527	1599.F02.gz43_212944
M00055610D:H09	ES 173	639255	1599.F09.gz43_213056
M00055611C:E03	ES 173	551896	1599.F24.gz43_213296
M00055612A:H05	ES 173	645031	1599.G13.gz43_213121
M00055615C:E01	ES 173	640868	1599.I08.gz43_213043
M00055615D:C07	ES 173	639703	1599.I11.gz43_213091
M00055617A:H12	ES 173	641254	1599.J09.gz43_213060
M00055619A;C03	ES 173	213631	1599.K12.gz43_213109
M00055619B:H04	ES 173	639480	1599.K19.gz43_213221
M00055619C:D06	ES 173	647518	1599.K24.gz43_213301
M00055619C:F07	ES 173	640221	1599.L01.gz43_212934
M00055619D:A04	ES 173	640382	1599.L05.gz43_212998

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Table 13			
CloneID	ES No	ClusterID	SequenceName
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M00055621B:G03	ES 173	561626	1599.L23.gz43_213286
M00055623D:G05	ES 173	641069	1599.N09.gz43_213064
M00055628B:B07	ES 173	607138	1599,P21,gz43_213258
M00055629A;D08	ES 173	647266	1600.A15.gz43_213531
M00055629B:G09	ES 173	455075	1600.A22.gz43_213643
M00055630B:E09	ES 173	561180	1600.B13.gz43_213500
M00055630B:G04	ES 173	645004	1600.B15.gz43 213532
M00055632A:B11	ES 173	559423	1600.B24.gz43_213676
M00055632D:A06	ES 173	640799	1600.C10.gz43_213453
M00055633B:A08	ES 173	634012	1600.C21.gz43 213629
M00055633B:G02	ES 173	131348	1600.C24.gz43 213677
M00055633D:A02	ES 173	649717	1600.D04.gz43 213358
M00055634C:C10	ES 173	446659	1600.D21.gz43_213630
M00055634C:F09	ES 173	166041	1600.D23.gz43 213662
M00055636A:H12	ES 173	650180	1600.E24.gz43_213679
M00055637B:A01	ES 173	642361	1600.F24.gz43_213680
M00055637B:H12	ES 173	641253	1600.G05.gz43_213377
M00055638A:A10	ES 173	554335	1600.G13.gz43 213505
M00055638D:D07	ES 173	639934	1600.H02.gz43_213330
M00055638D:E09	ES 173	646695	1600.H04.gz43_213362
M00055639D:D03	ES 173	548959	1600.I01.gz43_213315
M00055639D:F08	ES 173	640956	1600.I03.gz43 213347
M00055640A:G03	ES 173	635965	1600.I07.gz43_213411
M00055640B:C01	ES 173	472129	1600.I09.gz43_213443
M00055640C:E06	ES 173	557401	1600.I12.gz43 213491
M00055640C:F05	ES 173	467381	1600,I13.gz43_213507
M00055640C:F08	ES 173	151279	1600.I14.gz43_213523
M00055641A:C12	ES 173	562000	1600.I19.gz43_213603
M00055642D:A05	ES 173	553516	1600.J23.gz43_213668
M00055643A:C01	ES 173	451615	1600.K05.gz43_213381
M00055643B:E05	ES 173	642246	1600.K11.gz43_213477
M00055643C:G10	ES 173	645261	1600.K18.gz43_213589
M00055643D:A05	ES 173	656268	1600.K20.gz43_213621
M00055643D:G11	ES 173	561069	1600.K24.gz43 213685
M00055644A:D10	ES 173	549124	1600.L02.gz43 213334
M00055644A:D12	ES 173	554722	1600.L03.gz43_213350
M00055644B:H12	ES 173	379040	1600.L07.gz43_213414
M00055645A:C07	ES 173	156097	1600.L19.gz43_213606
M00055646C:B04	ES 173	646352	1600.M13.gz43_213511
M00055647A:H10	ES 173	446621	1600.M19.gz43_213607
M00055647B:A05	ES 173	641210	1600.M20.gz43 213623
M00055647C:D02	ES 173	639886	1600.N04.gz43 213368
M00055647D:B11	ES 173	467563	1600.N11.gz43_213480

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CloneID	ES No	ClusterID	SequenceName
M00055648B:C01	ES 173	453441	1600.N19.gz43_213608
M00055649A:H07	ES 173	640320	1600,O11.gz43_21348
M00055649C:F02	ES 173	491368	1600.O16.gz43_21356
M00055650A:B05	ES 173	553285	1600.O21.gz43_21364
M00055650C:F12	ES 173	640029	1600.P09.gz43_21345
M00055651A:D06	ES 173	638779	1600.P16.gz43_21356
M00055651A:E06	ES 173	641702	1600.P17.gz43_21357
M00055651B:F08	ES 173	458736	1600.P20.gz43_21362
M00055651C:E01	ES 173	550515	1600.P24.gz43_21369
M00055651C:F07	ES 173	643513	1669.A01.gz43_26068
M00055652A:G11	ES 173	554510	1669.A07.gz43_26078
M00055652B:B11	ES 173	618998	1669.A10.gz43 26083
M00055652B:F12	ES 173	406734	1669.A12.gz43 26086
M00055653A:G08	ES 173	649631	1669.A24.gz43 26105
M00055653B:C03	ES 173	647333	1669.B06.gz43 26076
M00055653B:E12	ES 173	86311	1669.B09.gz43 26081
M00055653B:G01	ES 173	644971	1669.B10.gz43 26083
M00055653B:G11	ES 173	640259	1669.B11.gz43 26084
M00055653D:H02	ES 173	463312	1669.B23.gz43 26104
M00055654A:B07	ES 173	462247	1669.C01.gz43 26068
M00055654B:G09	ES 173	472226	1669.C06.gz43 26076
M00055654C:A05	ES 173	641469	1669.C08.gz43_26080
M00055654C:C10	ES 173	638971	1669.C09.gz43 26081
M00055654C:D03	ES 173	556488	1669.C10.gz43_26083
M00055654D:F02	ES 173	650677	1669.C17.gz43 26094
M00055655A:A09	ES 173	642111	1669.C18.gz43_26096
M00055655B:B08	ES 173	642411	1669.C22.gz43_26102
M00055656A:E09	ES 173	643804	1669.D11.gz43_26085
M00055657A:B04	ES 173	450883	1669.E08.gz43_26080
M00055659A:A08	ES 173	648773	1669.F11.gz43_26085
M00055659C:B10	ES 173	639706	1669.F21.gz43_26101
M00055659C:D06	ES 173	639674	1669.F24.gz43_26106
M00055660A:A06	ES 173	452687	1669.G10.gz43_26083
M00055660A;C05	ES 173	503275	1669.G11.gz43_26085
M00055660A:C08	ES 173	643563	1669.G12.gz43_26086
M00055660B:H02	ES 173	589483	1669.G17.gz43_26094
M00055661B:E07	ES 173	455778	1669.H06.gz43_26077
M00055661C:E11	ES 173	658447	1669.H12.gz43_26087
M00055662B:F11	ES 173	447585	1669.I04.gz43_26074
M00055662C:C11	ES 173	523875	1669.I07.gz43_26079
M00055662C:D05	ES 173	645758	1669.I08.gz43_26080
M00055662C:D12	ES 173	641379	1669.I09.gz43_26082
M00055662C:H06	ES 173	446188	1669.I12.gz43_26087
M00055662D:A09	ES 173	639596	1669.I15.gz43 26091

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Table 15			
CloneID	ES No	ClusterID	SequenceName
M00055662D:B07	ES 173	639665	1669.I17.gz43_260951
M00055662D:E05	ES 173	640018	1669.I20.gz43_260999
M00055662D:F09	ES 173	559575	1669.I21.gz43_261015
M00055663A:B02	ES 173	641091	1669.I22.gz43_261031
M00055663A;G04	ES 173	640261	1669.I24.gz43_261063
M00055663A:H10	ES 173	641658	1669.J01.gz43_260696
M00055663B:E02	ES 173	640078	1669.J04.gz43_260744
M00055663C:D05	ES 173	658271	1669.J07.gz43_260792
M00055663C:F05	ES 173	512521	1669.J08.gz43_260808
M00055663D:B05	ES 173	643909	1669.J10.gz43_260840
M00055664B:G08	ES 173	640211	1669,J16.gz43_260936
M00055665B:A12	ES 173	451906	1669.K06.gz43_260777
M00055665B:B10	ES 174	465104	1669.K07.gz43_260793
M00055665D:F11	ES 174	640179	1669.K18.gz43_260969
M00055666A:G04	ES 174	642078	1669.K23.gz43_261049
M00055666B:C03	ES 174	448484	1669.L07.gz43_260794
M00055666B:E06	ES 174	640678	1669.L11.gz43_260858
M00055666C;C11	ES 174	639804	1669.L12.gz43_260874
M00055666D:D08	ES 174	639932	1669.L16.gz43_260938
M00055667A:B12	ES 174	642332	1669.L20.gz43_261002
M00055667A:H10	ES 174	456561	1669.L23.gz43_261050
M00055667B:C08	ES 174	639752	1669.M01.gz43_260699
M00055667C:F07	ES 174	646894	1669.M10.gz43_260843
M00055667D:B01	ES 174	561894	1669.M12.gz43_260875
M00055668B:A10	ES 174	550999	1669.M19.gz43_260987
M00055668B:B07	ES 174	557401	1669.M20.gz43_261003
M00055668B:D05	ES 174	639906	1669.M21.gz43_261019
M00055668C:A04	ES 174	642068	1669.N02.gz43_260716
M00055668C:F05	ES 174	648340	1669.N05.gz43_260764
M00055668D:E11	ES 174	556750	1669.N10.gz43_260844
M00055669B:G02	ES 174	648352	1669.N18.gz43_260972
M00055669D:B08	ES 174	639743	1669,O02.gz43_260717
M00055670A:B04	ES 174	648352	1669.007.gz43_260797
M00055670D:F02	ES 174	641683	1669.O17.gz43_260957
M00055671A:H03	ES 174	518569	1669.P01.gz43_260702
M00055671B:B02	ES 174	642230	1669.P03.gz43_260734
M00055672D:F06	ES 174	640092	1670.A08.gz43_261183
M00055673A:E09	ES 174	95617	1670.A11.gz43 261231
M00055673B:A04	ES 174	645146	1670.A16.gz43_261311
M00055673B:B08	ES 174	639469	1670.A18.gz43_261343
M00055673D:C01	ES 174	639805	1670.B07.gz43_261168
M00055674A:C11	ES 174	651088	1670.B12.gz43_261248
M00055674C:E05	ES 174	645968	1670.B21.gz43_261392

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055675A:G03	ES 174	643522	1670.C06.gz43_261153
M00055675D:E04	ES 174	650673	1670.C14.gz43_261281
M00055676B:D05	ES 174	446242	1670.C20.gz43_261377
M00055676D:H07	ES 174	426366	1670.D05.gz43_261138
M00055678C:H05	ES 174	289328	1670.D14.gz43_261282
M00055679B:E03	ES 174	642321	1670.D23.gz43_261426
M00055680B:D09	ES 174	553642	1670.E15.gz43_261299
M00055680B:H04	ES 174	647427	1670.E17.gz43_261331
M00055680C;E01	ES 174	493622	1670.E20.gz43_261379
M00055681A:F02	ES 174	416808	1670.E24.gz43_261443
M00055681D:F11	ES 174	642502	1670.F18.gz43_261348
M00055682A:B07	ES 174	637966	1670.F22.gz43_261412
M00055682C:D06	ES 174	641618	1670.G09.gz43_261205
M00055682D:G02	ES 174	641902	1670,G19.gz43_261365
M00055683B:D10	ES 174	640645	1670,H06.gz43_261158
M00055683C:A03	ES 174	557625	1670.H10.gz43_261222
M00055683C:D11	ES 174	644699	1670.H14.gz43_261286
M00055683C:H11	ES 174	553921	1670.H17.gz43_261334
M00055684B:D06	ES 174	599838	1670.I10.gz43_261223
M00055684D:A10	ES 174	506901	1670.I18.gz43_261351
M00055685A:E10	ES 174	601137	1670.I23.gz43_261431
M00055685A:F02	ES 174	647856	1670.I24.gz43_261447
M00055685A:F06	ES 174	447802	1670.J02.gz43_261096
M00055685B:E12	ES 174	645909	1670.J06.gz43_261160
M00055685D:B06	ES 174	649558	1670.J18.gz43_261352
M00055686B:D03	ES 174	641624	1670.K06.gz43_261161
M00055686D:E04	ES 174	641726	1670.K12.gz43_261257
M00055686D:E10	ES 174	641728	1670.K13.gz43_261273
M00055687C:B04	ES 174	554564	1670.K24.gz43_261449
M00055687C:B11	ES 174	649284	1670.L01.gz43_261082
M00055687C:F01	ES 174	643594	1670.L05.gz43_261146
M00055687C:F07	ES 174	639136	1670.L06.gz43_261162
M00055688A:A02	ES 174	641287	1670.L09.gz43_261210
M00055688A:E04	ES 174	649921	1670.L12.gz43_261258
M00055689B;F04	ES 174	446675	1670.M05.gz43_261147
M00055689C:B03	ES 174	634122	1670.M08.gz43_261195
M00055689C:C03	ES 174	634122	1670,M10.gz43_261227
M00055689D:G01	ES 174	567636	1670.M18.gz43_261355
M00055691A:D08	ES 174	640826	1670.M22.gz43_261419
M00055691B:E07	ES 174	640889	1670,N03.gz43_261116
M00055691C:E02	ES 174	646638	1670.N06.gz43_261164
M00055691D:B07	ES 174	642644	1670.N11.gz43_261244
M00055691D:E04	ES 174	502609	1670.N13.gz43_261276
M00055692A:E05	ES 174	561920	1670.N18.gz43_261356

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CloneID	ES No	ClusterID	SequenceName
M00055695D:E01	ES 174	645431	1670.P12.gz43_261262
M00055699C:D02	ES 174	41141	1671.B11.gz43_261616
M00055700B:E10	ES 174	553985	1671.B23.gz43_261808
M00055701A:B04	ES 174	560546	1671.C12.gz43_261633
M00055701B:C03	ES 174	642039	1671.C17.gz43_261713
M00055702C:D01	ES 174	646028	1671.D09.gz43_261586
M00055702C:F11	ES 174	640947	1671.D10.gz43_261602
M00055702D:H10	ES 174	605761	1671.D11.gz43_261618
M00055704A:D09	ES 174	642062	1671.E08.gz43_261571
M00055704B:C04	ES 174	551415	1671.E12.gz43_261635
M00055705C:F12	ES 174	471268	1671.F17.gz43_261716
M00055705C:G01	ES 174	641876	1671.F18.gz43_261732
M00055706B:H12	ES 174	463548	1671.G10.gz43_261605
M00055707A:B07	ES 174	649429	1671.G14.gz43_261669
M00055707A:E07	ES 174	468316	1671.G17.gz43_261717
M00055707A:F11	ES 174	453006	1671.G20.gz43_261765
M00055707B:E02	ES 174	640879	1671.G22.gz43_261797
M00055709D:G10	ES 174	417259	1671.H17.gz43_261718
M00055710B:E04	ES 174	641700	1671.H22.gz43_261798
M00055711B:B08	ES 174	561422	1671.I07.gz43_261559
M00055711C:A07	ES 174	463368	1671.I11.gz43_261623
M00055711D:H12	ES 174	558890	1671.I21.gz43_261783
M00055713C:B06	ES 174	455552	1671.J08.gz43_261576
M00055713C:D11	ES 174	469688	1671.J10.gz43_261608
M00055713C:F12	ES 174	558086	1671.J11.gz43_261624
M00055713C:H01	ES 174	446933	1671.J12.gz43_261640
M00055715A:D10	ES 174	463368	1671.K08.gz43_261577
M00055715C:C11	ES 174	640534	1671.K20.gz43_261769
M00055717A:C06	ES 174	644325	1671.L10.gz43_261610
M00055717A:H01	ES 174	521888	1671.L12.gz43_261642
M00055717B:A03	ES 174	451401	1671.L14.gz43_261674
M00055717B:E04	ES 174	421826	1671.L17.gz43_261722
M00055717C:B07	ES 174	536415	1671.L23.gz43_261818
M00055718A:F01	ES 174	478192	1671.M09.gz43_261595
M00055718A:H05	ES 174	640282	1671.M10.gz43_261611
M00055718B:H11	ES 174	465589	1671.M17.gz43_261723
M00055719A:A06	ES 174	646105	1671.N09.gz43_261596
M00055719A:D11	ES 174	557710	1671.N12.gz43_261644
M00055719A:G04	ES 174	645505	1671.N17.gz43_261724
M00055719A:G12	ES 174	510195	1671.N18.gz43_261740
M00055719C:A09	ES 174	640282	1671.N23.gz43_261820
M00055720B:D06	ES 174	645344	1671.O12.gz43_261645
M00055721A:A07	ES 174	466265	1671.O22.gz43_261805
M00055721B:D08	ES 174	642288	1671.P07.gz43_261566

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055722A:B05	ES 174	492094	1671.P18.gz43_261742
M00055722A:C04	ES 174	600115	1671.P21.gz43_261790
M00055722B:A01	ES 174	498194	1672.A01.gz43_261841
M00055722B:G12	ES 174	645171	1672.A06.gz43_261921
M00055722C:E11	ES 174	650303	1672.A12.gz43_262017
M00055722C:F11	ES 174	642491	1672.A13.gz43_262033
M00055722D:A07	ES 174	641056	1672.A14.gz43_262049
M00055722D:B10	ES 174	460967	1672.A17.gz43 262097
M00055722D:E05	ES 174	544797	1672.A19.gz43_262129
M00055722D:G12	ES 174	470769	1672.A23.gz43_262193
M00055723A:B08	ES 174	556326	1672.B03.gz43_261874
M00055723A:C04	ES 174	647688	1672.B04.gz43 261890
M00055723A:F04	ES 174	649360	1672.B07.gz43_261938
M00055723B:A09	ES 174	642078	1672.B11.gz43_262002
M00055723B:C03	ES 174	642197	1672.B14.gz43_262050
M00055723B;H08	ES 174	46976	1672.B17.gz43_262098
M00055723C:A08	ES 174	648472	1672.B18.gz43_262114
M00055723C:B02	ES 174	489426	1672.B19.gz43_262130
M00055723C:F09	ES 174	503923	1672.B22.gz43_262178
M00055723D:C04	ES 174	522497	1672.C03.gz43_261875
M00055723D:E05	ES 174	649106	1672.C06.gz43_261923
M00055724A:C12	ES 174	284586	1672.C12.gz43_262019
M00055724A:E03	ES 174	645018	1672,C15.gz43_262067
M00055724A:G08	ES 174	504568	1672.C17.gz43_262099
M00055724B:D04	ES 174	467822	1672.C20.gz43_262147
M00055724B:G03	ES 174	642722	1672.C23.gz43_262195
M00055724B:H03	ES 174	557221	1672.D02.gz43_261860
M00055724D;A02	ES 174	559828	1672.D17.gz43_262100
M00055725A:F12	ES 174	641305	1672.E02.gz43_261861
M00055725A:G07	ES 174	557833	1672.E03.gz43_261877
M00055725D:B02	ES 174	522548	1672.E14.gz43_262053
M00055725D:F05	ES 174	452212	1672.E19.gz43_262133
M00055726C:F01	ES 174	645004	1672.F16.gz43_262086
M00055726D:H12	ES 174	489249	1672.F24.gz43_262214
M00055727A:G01	ES 174	507188	1672.G02.gz43_261863
M00055727B:F10	ES 174	642535	1672.G12.gz43_262023
M00055727C:B02	ES 174	559776	1672.G16.gz43_262087
M00055727D:G01	ES 174	641070	1672.H02.gz43_261864
M00055727D:H04	ES 174	639711	1672.H04.gz43_261896
M00055728B:G11	ES 174	647060	1672.H16.gz43_262088
M00055728C:B08	ES 174	455113	1672.H17.gz43_262104
M00055728D:F02	ES 174	554708	1672.H22.gz43_262184
M00055729A:B12	ES 174	672032	1672.I02.gz43_261865
M00055729B:G03	ES 174	641101	1672.I12.gz43_262025

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CloneID	ES No	ClusterID	SequenceName
M00055729C:D11	ES 174	642315	1672.I16.gz43 262089
M00055729C:E02	ES 174	477064	1672.I17.gz43_262105
M00055729D:A06	ES 174	546705	1672.I18.gz43_262121
M00055729D:F05	ES 174	644609	1672.I21.gz43 262169
M00055731A:C03	ES 174	461583	1672,J13.gz43 262042
M00055731A:G05	ES 174	640704	1672.J17.gz43_262106
M00055731A:H07	ES 174	235397	1672.J18.gz43_262122
M00055731C:F09	ES 174	454383	1672.K06.gz43_261931
M00055731C:H07	ES 174	639458	1672.K08.gz43_261963
M00055731D:B10	ES 174	646129	1672.K11.gz43_262011
M00055731D:E12	ES 174	555681	1672.K11.gz43_262041
M00055731D:F09	ES 174	640879	1672.K15.gz43_262075
M00055732A:B04	ES 174	651085	1672.K18.gz43_262123
M00055732A:C12	ES 174	322255	1672.K19.gz43_262139
M00055732A:F09	ES 174	642557	1672.K21.gz43_262171
M00055733A:G04	ES 174	642631	1672.L11.gz43_262012
M00055733B:F11	ES 174	504501	1672.L16.gz43_262092
M00055733B:H05	ES 174	419443	1672.L17.gz43 262108
M00055733C:G02	ES 174	383609	1672.L18.gz43_262124
M00055733C:H12	ES 174	650397	1672.L20.gz43_262156
M00055733D:B11	ES 174	561968	1672.L21.gz43 262172
M00055733D:D04	ES 174	645151	1672.L23.gz43 262204
M00055733D:H01	ES 174	416377	1672.M04.gz43 261901
M00055734D:A02	ES 174	461474	1672.M16.gz43_262093
M00055734D:D01	ES 174	646609	1672.M19.gz43 262141
M00055734D:E05	ES 174	642376	1672.M21.gz43 262173
M00055734D:E10	ES 174	645165	1672.M22.gz43 262189
M00055734D:G12	ES 174	558146	1672.M23.gz43 262205
M00055735A:B05	ES 174	647669	1672,N01,gz43 261854
M00055735B:F11	ES 175	642111	1672.N10.gz43 261998
M00055735C:G08	ES 175	648667	1672.N14.gz43_262062
M00055735D:C01	ES 175	502826	1672,N18,gz43 262126
M00055735D:E10	ES 175	556336	1672.N19.gz43 262142
M00055736A:B11	ES 175	648265	1672.N23,gz43 262206
M00055736B:C08	ES 175	643897	1672.004.gz43 261903
M00055736C:D06	ES 175	466434	1672.Q09.gz43 261983
M00055736D:D11	ES 175	642263	1672.O15.gz43 262079
M00055737B:A03	ES 175	640266	1672.020.gz43_262159
M00055738B:H11	ES 175	100821	1672.P13.gz43_262048
M00055738C:A12	ES 175	647556	1672.P14.gz43_262064
M00055738C:C02	ES 175	647906	1672.P15.gz43_262080
M00055738D:G08	ES 175	645085	1672.P20.gz43_262160
M00055739A:B02	ES 175	550049	1672.P21.gz43 262176

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Labic 15			
CloneID	ES No	ClusterID	SequenceName
M00055739A:B04	ES 175	608540	1672.P22.gz43_262192
M00055739A:C09	ES 175	558301	1681.A01.gz43_296936
M00055739B:B06	ES 175	468613	1681.A03.gz43_296968
M00055739B:D09	ES 175	644836	1681.A04.gz43_296984
M00055739B:H08	ES 175	568467	1681.A08.gz43_297048
M00055739C:D11	ES 175	648481	1681.A10.gz43_297080
M00055739D:B12	ES 175	644684	1681.A13.gz43_297128
M00055739D:C03	ES 175	463815	1681.A14.gz43_297144
M00055739D:E04	ES 175	644919	1681.A16.gz43_297176
M00055740A:B03	ES 175	462245	1681.A19.gz43_297224
M00055740B:C06	ES 175	643142	1681.A21.gz43_297256
M00055740B:F09	ES 175	463824	1681.A24.gz43_297304
M00055740C;A07	ES 175	648905	1681.B03.gz43_296969
M00055740C:E06	ES 175	644937	1681.B04.gz43_296985
M00055740D:G12	ES 175	150839	1681.B10.gz43_297081
M00055741A:D09	ES 175	640504	1681.B13.gz43_297129
M00055741B:B12	ES 175	640997	1681.B15.gz43_297161
M00055741C:A09	ES 175	526459	1681.B18.gz43_297209
M00055741D:H01	ES 175	548920	1681.C01.gz43_296938
M00055742B:H06	ES 175	645147	1681.C08.gz43_297050
M00055742C:A07	ES 175	640306	1681.C10.gz43_297082
M00055742C:C01	ES 175	419479	1681.C11.gz43_297098
M00055742D:H03	ES 175	413621	1681,C16.gz43_297178
M00055743B:E01	ES 175	389377	1681.C24.gz43_297306
M00055743C:C01	ES 175	508755	1681.D02.gz43_296955
M00055743C:D12	ES 175	463951	1681.D05.gz43_297003
M00055743D:D04	ES 175	649427	. 1681.D09.gz43_297067
M00055743D:E07	ES 175	448703	1681.D10.gz43_297083
M00055744A:B04	ES 175	642263	1681.D15.gz43_297163
M00055744B:B02	ES 175	640017	1681.D19.gz43_297227
M00055744B:C08	ES 175	218416	1681.D20.gz43_297243
M00055744C:D02	ES 175	642644	1681.E02.gz43_296956
M00055744D:A11	ES 175	447936	1681.E08.gz43_297052
M00055744D:F03	ES 175	645000	1681.E11.gz43_297100
M00055745A;H02	ES 175	453726	1681.E19.gz43_297228
M00055745B:H02	ES 175	470667	1681.E23.gz43_297292
M00055745C:A06	ES 175	468959	1681.E24.gz43_297308
M00055745C:G06	ES 175	552085	1681.F03.gz43_296973
M00055745D:F11	ES 175	556654	1681.F09.gz43_297069
M00055746A:C09	ES 175	644053	1681.F10.gz43_297085
M00055746C:F06	ES 175	587696	1681.F17.gz43_297197
M00055746C:F10	ES 175	471277	1681.F18.gz43_297213
M00055746C:G06	ES 175	644468	1681.F19.gz43_297229
M00055747C:E09	ES 175	529742	1681.Gl1.gz43_297102

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055748A:B03	ES 175	593343	1681.G19.gz43_297230
M00055748A:D07	ES 175	448251	1681,G21.gz43_297262
M00055748C:C07	ES 175	446925	1681.H03.gz43_296975
M00055748D:C03	ES 175	594040	1681.H09.gz43_297071
M00055749B:C10	ES 175	461917	1681.H17.gz43_297199
M00055749B:D12	ES 175	452504	1681.H18.gz43_297215
M00055749C:B03	ES 175	644105	1681.H19.gz43_297231
M00055749C:C04	ES 175	489040	1681.H20.gz43_297247
M00055749D:B07	ES 175	426366	1681.I01.gz43_296944
M00055749D:C01	ES 175	644105	1681.I02.gz43_296960
M00055749D:D06	ES 175	649356	1681,I03.gz43_296976
M00055749D:F12	ES 175	644342	1681.I04.gz43_296992
M00055750B:H01	ES 175	644510	1681.I17.gz43_297200
M00055750C:H10	ES 175	638943	1681.J01.gz43_296945
M00055750D:H06	ES 175	649763	1681.J05.gz43_297009
M00055751A:F06	ES 175	488432	1681.J09.gz43_297073
M00055752A:E10	ES 175	649702	1681.J20.gz43_297249
M00055752A:G10	ES 175	644479	1681.J23.gz43_297297
M00055752C:C06	ES 175	510724	1681.K03.gz43 296978
M00055752C:F06	ES 175	644345	1681.K05.gz43_297010
M00055752C:H07	ES 175	611927	1681.K07.gz43_297042
M00055752D:C01	ES 175	641144	1681.K09.gz43_297074
M00055752D:F01	ES 175	644342	1681.K10.gz43_297090
M00055753A:D04	ES 175	639703	1681.K16.gz43_297186
M00055753B:A02	ES 175	483042	1681.K18.gz43_297218
M00055753B:A06	ES 175	447485	1681.K19.gz43_297234
M00055753B:F10	ES 175	449438	1681.K22.gz43_297282
M00055753D:C06	ES 175	446676	1681.L03.gz43_296979
M00055754A;E04	ES 175	446355	1681.L08.gz43_297059
M00055754A:E07	ES 175	550223	1681.L09.gz43_297075
M00055754A:H06	ES 175	478833	1681.L11.gz43_297107
M00055755A:B11	ES 175	514838	1681.L18.gz43_297219
M00055755C:F12	ES 175	464171	1681.M08.gz43_297060
M00055755D:C09	ES 175	491544	1681.M09.gz43_297076
M00055756C:C03	ES 175	639750	1681.M16.gz43_297188
M00055756D;B05	ES 175	643941	1681.M20.gz43_297252
M00055756D:E05	ES 175	108479	1681.M21.gz43_297268
M00055757A;A07	ES 175	553675	1681.M24.gz43_297316
M00055757B;B07	ES 175	646552	1681.N06.gz43_297029
M00055757B;C04	ES 175	639420	1681.N07.gz43_297045
M00055757B:D06	ES 175	235194	1681.N09.gz43_297077
M00055757D;B05	ES 175	643984	1681.N18.gz43_297221
M00055758A:G02	ES 175	644461	1681.N21.gz43_297269
M00055758C:H10	ES 175	645603	1681.Q01.gz43 296950

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055758D:E02	ES 175	640264	1681.004.gz43_296998
M00055759A:B02	ES 175	492917	1681.008.gz43_297062
M00055759A:E01	ES 175	295143	1681.O11.gz43_297110
M00055759C:C07	ES 175	453188	1681.O18.gz43_297222
M00055760A:E06	ES 175	453324	1681.O23.gz43_297302
M00055761A:E06	ES 175	451508	1681.P15.gz43_297175
M00055761B:F01	ES 175	646453	1681.P21.gz43_297271
M00055761B:F09	ES 175	641968	1681.P22.gz43_297287
M00055761D:C03	ES 175	648320	1682.A06.gz43_262305
M00055761D:H08	ES 175	488680	1682.A09.gz43_262353
M00055762B:B11	ES 175	639981	1682.A14.gz43_262433
M00055762C:B04	ES 175	643940	1682.A16.gz43_262465
M00055762C:E06	ES 175	494378	1682.A17.gz43_262481
M00055762C:H07	ES 175	466147	1682.A20.gz43_262529
M00055763A:H02	ES 175	649360	1682.B03.gz43_262258
M00055763B:D07	ES 175	550562	1682.B05.gz43_262290
M00055763B:E09	ES 175	644278	1682.B08.gz43_262338
M00055763C:A10	ES 175	462815	1682.B12.gz43_262402
M00055763C:G08	ES 175	642564	1682.B14.gz43_262434
M00055763D:B11	ES 175	562870	1682.B15.gz43_262450
M00055764A:F02	ES 175	621636	1682.B18.gz43_262498
M00055764B:F05	ES 175	462393	1682.B22.gz43_262562
M00055764C:C07	ES 175	644765	1682.B24.gz43_262594
M00055765A:A04	ES 175	524470	1682.C10.gz43_262371
M00055765A:B02	ES 175	584179	1682.C11.gz43_262387
M00055765A;C11	ES 175	642204	1682.C16.gz43_262467
M00055765A:G02	ES 175	629002	1682.C20.gz43_262531
M00055765D:E06	ES 175	642460	1682.D11.gz43_262388
M00055765D:F04	ES 175	644853	1682.D13.gz43_262420
M00055766A:A01	ES 175	462742	1682.D15.gz43_262452
M00055766A:D08	ES 175	641576	1682.D17.gz43_262484
M00055766A:E08	ES 175	639232	1682.D18.gz43_262500
M00055766A:H03	ES 175	558719	1682.D20.gz43_262532
M00055766B:B01	ES 175	447224	1682.D23.gz43_262580
M00055766B:E11	ES 175	637387	1682.E04.gz43_262277
M00055766B:H10	ES 175	642852	1682.E08.gz43_262341
M00055766C:C07	ES 175	464091	1682.E10.gz43_262373
M00055766C:C11	ES 175	448944	1682.E11.gz43_262389
M00055766C:E05	ES 175	447712	1682.E12.gz43_262405
M00055766C:G12	ES 175	451037	1682.E15.gz43_262453
M00055766D:C05	ES 175	562021	1682.E18.gz43_262501
M00055766D:D05	ES 175	50351	1682.E20.gz43_262533
M00055766D:F09	ES 175	560393	1682.E24.gz43_262597
M00055767A:D10	ES 175	634660	1682.F04.gz43_262278

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Table 15			
CloneID	ES No	ClusterID	SequenceName
M00055767B:D02	ES 175	642082	1682.F11.gz43_262390
M00055767D:A12	ES 175	642082	1682.F17.gz43_262486
M00055767D:E07	ES 175	642417	1682.F20.gz43_262534
M00055767D:F04	ES 175	546740	1682.F21.gz43_262550
M00055768A:B05	ES 175	648996	1682.F24.gz43_262598
M00055768B:H12	ES 175	541499	1682.G13.gz43 262423
M00055768D:G02	ES 175	555140	1682.G22.gz43 262567
M00055770A:F01	ES 175	640997	1682.G24.gz43_262599
M00055770A:G08	ES 175	644173	1682,H02.gz43_262248
M00055770A:H11	ES 175	530238	1682.H03.gz43_262264
M00055770B:D06	ES 175	446757	1682.H06.gz43_262312
M00055770B:F06	ES 175	642474	1682.H07.gz43_262328
M00055770C:A02	ES 175	551167	1682.H08.gz43_262344
M00055770C:D01	ES 175	234606	1682.H09.gz43_262360
M00055770C:H11	ES 175	562876	1682.H12.gz43_262408
M00055770D:E10	ES 175	464510	1682.H19.gz43_262520
M00055771A:A11	ES 175	557797	1682.H22.gz43_262568
M00055771B:G05	ES 175	471268	1682.I10.gz43_262377
M00055771C:A11	ES 175	453079	1682.I12.gz43_262409
M00055771C:D09	ES 175	509410	1682.I13.gz43_262425
M00055771C:F05	ES 175	642604	1682.I15.gz43_262457
M00055772A:C10	ES 175	642036	1682.I24.gz43_262601
M00055772A:E12	ES 175	648588	1682.J01.gz43_262234
M00055772A:H08	ES 175	642791	1682.J04.gz43_262282
M00055772C:B09	ES 175	452503	1682.J08.gz43_262346
M00055772C:E11	ES 175	463951	1682.J10.gz43_262378
M00055772C:G08	ES 175	138470	1682.J12.gz43_262410
M00055772D:C10	ES 175	641563	1682.J15.gz43_262458
M00055772D:D03	ES 175	517912	1682.J16.gz43_262474
M00055772D:F10	ES 175	450553	1682.J18.gz43_262506
M00055772D:F11	ES 175	642558	1682.J19.gz43_262522
M00055772D:H04	ES 175	466971	1682.J20.gz43_262538
M00055773A:F05	ES 175	644451	1682.J22.gz43_262570
M00055773B:A07	ES 175	644914	1682.J24.gz43_262602
M00055773C:C09	ES 175	446163	1682.K04.gz43_262283
M00055773C:D12	ES 175	650272	1682.K05.gz43_262299
M00055773C:H12	ES 175	454540	1682.K06.gz43_262315
M00055773D:G11	ES 175	642637	1682.K09.gz43_262363
M00055774A:D04	ES 175	449975	1682.K14.gz43_262443
M00055774B:F07	ES 175	647940	1682.K18.gz43_262507
M00055774B:H01	ES 175	641185	1682.K20.gz43_262539
M00055774C:D09	ES 175	640617	1682.K22.gz43_262571
M00055774D:A05	ES 175	642417	1682.L02.gz43_262252

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055774D:B07	ES 176	553380	1682.L03.gz43_262268
M00055774D:E02	ES 176	458979	1682,L06.gz43_262316
M00055775A:G09	ES 176	101499	1682.L12.gz43_262412
M00055775B:A06	ES 176	447206	1682.L16.gz43_262476
M00055775B;G02	ES 176	644240	1682.L20.gz43_262540
M00055775B:G04	ES 176	559662	1682.L21.gz43_262556
M00055775C:B02	ES 176	645327	1682.L24.gz43_262604
M00055775C:B10	ES 176	553877	1682.M02.gz43_262253
M00055775C:D08	ES 176	642265	1682.M03.gz43_262269
M00055775C;E10	ES 176	641056	1682.M04.gz43_262285
M00055778A:E09	ES 176	621635	1682.M16.gz43_262477
M00055778A:F09	ES 176	461835	1682.M18.gz43_262509
M00055778A:F12	ES 176	642564	1682.M19.gz43_262525
M00055778A:G02	ES 176	509027	1682.M20.gz43_262541
M00055778B:C03	ES 176	645036	1682.M22.gz43_262573
M00055778B:E01	ES 176	458425	1682.M24.gz43_262605
M00055778C:D09	ES 176	494767	1682.N06.gz43_262318
M00055778D:A03	ES 176	642095	1682.N11.gz43_262398
M00055778D:C03	ES 176	639829	1682.N12.gz43_262414
M00055779A:B06	ES 176	642166	1682.N15.gz43_262462
M00055779A:H05	ES 176	480035	1682.N20.gz43_262542
M00055779B:F06	ES 176	556476	1682.N24.gz43_262606
M00055780A:C04	ES 176	89082	1682.O13.gz43_262431
M00055780A:C06	ES 176	561558	1682.O14.gz43_262447
M00055780A:E01	ES 176	650451	1682.O15.gz43_262463
M00055780A:E03	ES 176	642491	1682.O16.gz43_262479
M00055780A:E11	ES 176	450211	1682.O17.gz43_262495
M00055780A:F07	ES 176	641496	1682.O18.gz43_262511
M00055780A:G04	ES 176	453090	1682.O20.gz43_262543
M00055780C:E02	ES 176	641174	1682.P06.gz43_262320
M00055780C:E10	ES 176	502614	1682,P07.gz43_262336
M00055780D:D09	ES 176	642332	1682.P12.gz43_262416
M00055780D:F08	ES 176	453091	1682.P13.gz43_262432
M00055781A:B04	ES 176	642147	1682.P19.gz43_262528
M00055782B:C08	ES 176	643230	1683.A19.gz43_262897
M00055782C:H01	ES 176	459064	1683.B06.gz43_262690
M00055782D:H01	ES 176	649883	1683.B11.gz43_262770
M00055783A:C06	ES 176	642184	1683.B14.gz43_262818
M00055783A:F03	ES 176	644486	1683.B18.gz43_262882
M00055783B:A04	ES 176	523590	1683.B20.gz43_262914
M00055783C:A05	ES 176	506285	1683.C01.gz43_262611
M00055783D:B02	ES 176	497971	1683.C05.gz43_262675
M00055784A:B07	ES 176	140909	1683.C12.gz43_262787
M00055784A:D05	ES 176	456626	1683.C16.gz43_262851

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Table 13	T 70.31		
CloneID	ES No	ClusterID	SequenceName
M00055784A:G10	ES 176	450384	1683.C19.gz43_262899
M00055784C:H02	ES 176	675768	1683.D03.gz43_262644
M00055784D:E03	ES 176	457847	1683.D05.gz43_262676
M00055785A:H08	ES 176	549434	1683.D09.gz43_262740
M00055785B:B06	ES 176	650076	1683.D12.gz43_262788
M00055785D:C06	ES 176	672601	1683.E01.gz43_262613
M00055786D:E10	ES 176	461990	1683.E19.gz43_262901
M00055787B:E02	ES 176	541209	1683.F03.gz43_262646
M00055787B:F10	ES 176	648905	1683.F04.gz43_262662
M00055787C:D09	ES 176	643350	1683.F10.gz43_262758
M00055787C:E12	ES 176	449836	1683.F12.gz43_262790
M00055787D:B07	ES 176	642318	1683.F16.gz43_262854
M00055789C:C12	ES 176	640799	1683.G23.gz43_262967
M00055789C:F10	ES 176	463290	1683.H01.gz43_262616
M00055789D:B04	ES 176	642146	1683.H03.gz43_262648
M00055789D:C06	ES 176	621635	1683.H04.gz43_262664
M00055790B:A08	ES 176	638962	1683.H14.gz43_262824
M00055790B:D05	ES 176	553546	1683.H18.gz43_262888
M00055790C:C02	ES 176	415950	1683.H22.gz43_262952
M00055790D:G10	ES 176	467306	1683.I04.gz43_262665
M00055791B:E02	ES 176	510545	1683.I11.gz43_262777
M00055791C:A02	ES 176	396191	1683.I15.gz43_262841
M00055792B:D08	ES 176	396149	1683.J14.gz43_262826
M00055793A:H09	ES 176	554117	1683.K06.gz43_262699
M00055793B:B06	ES 176	641919	1683.K08.gz43_262731
M00055794A:D08	ES 176	605761	1683.L08.gz43_262732
M00055794B:F04	ES 176	561718	1683.L17.gz43_262876
M00055794C:D10	ES 176	457396	1683.L21.gz43_262940
M00055795A:F09	ES 176	446595	1683,M09.gz43_262749
M00055795A:F12	ES 176	454485	1683.M10.gz43_262765
M00055795B;F09	ES 176	641700	1683,M15.gz43_262845
M00055795C:B10	ES 176	388085	1683,M20.gz43_262925
M00055795D:E09	ES 176	645508	1683,N05.gz43_262686
M00055795D:F08	ES 176	450218	1683.N06.gz43_262702
M00055795D:H08	ES 176	77144	1683.N11.gz43 262782
M00055796A:A08	ES 176	437580	1683.N15.gz43 262846
M00055796B:G05	ES 176	644047	1683,O05.gz43 262687
M00055796C:E11	ES 176	647639	1683,O10.gz43 262767
M00055796D:E06	ES 176	553979	1683.O16.gz43_262863
M00055796D:E10	ES 176	640525	1683.O17.gz43 262879
M00055797B:A11	ES 176	417617	1683.O23.gz43_262975
M00055797B:B04	ES 176	639807	1683.024.gz43_262991
M00055797C:F08	ES 176	517224	1683,P09.gz43_262752
M00055798B:D12	ES 176	418340	1684.A04.gz43_263041

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CloneID	ES No	ClusterID	SequenceName
M00055798B:F02	ES 176	644915	1684.A05.gz43_263057
M00055798B:G04	ES 176	645560	1684.A06.gz43_263073
M00055798D:A10	ES 176	455028	1684.A10.gz43_263137
M00055798D:H10	ES 176	483726	1684.A13.gz43_263185
M00055799B:G02	ES 176	674574	1684.A23.gz43_263345
M00055799C:D09	ES 176	645700	1684.A24.gz43_263361
M00055799D:F11	ES 176	640334	1684.B06.gz43_263074
M00055800A:F01	ES 176	462247	1684.B14.gz43_263202
M00055800A:F02	ES 176	490414	1684.B15.gz43_263218
M00055800C:E11	ES 176	640537	1684.B22.gz43_263330
M00055800D:B03	ES 176	462249	1684.C04.gz43_263043
M00055801D:E06	ES 176	564382	1684.C23.gz43_263347
M00055802A:C03	ES 176	494767	1684.D06.gz43_263076
M00055802B:H03	ES 176	486134	1684.D17.gz43_263252
M00055802C:E12	ES 176	643991	1684.D22.gz43_263332
M00055803A:C06	ES 176	513238	1684.E06.gz43_263077
M00055803A:F03	ES 176	455405	1684.E10.gz43_263141
M00055803A:G08	ES 176	642653	1684.E11.gz43_263157
M00055803B:A11	ES 176	530774	1684.E13.gz43_263189
M00055803B:E10	ES 176	557344	1684.E18.gz43_263269
M00055804B:C02	ES 176	639194	1684.F18.gz43_263270
M00055804D:F02	ES 176	463821	1684.G01.gz43_262999
M00055805A:A02	ES 176	640222	1684.G05.gz43_263063
M00055805A:C11	ES 176	641059	1684.G06.gz43_263079
M00055805B:C08	ES 176	642198	1684.G12.gz43_263175
M00055805C:D10	ES 176	630259	1684.G17.gz43_263255
M00055805D:C11	ES 176	640617	1684.G21.gz43_263319
M00055805D:H01	ES 176	529325	1684.G23.gz43_263351
M00055806B:B10	ES 176	644314	1684.H08.gz43_263112
M00055806B:D04	ES 176	464040	1684,H10.gz43_263144
M00055806C:D07	ES 176	496051	1684.H13.gz43_263192
M00055806C:E09	ES 176	375814	1684.H14.gz43_263208
M00055806C;G01	ES 176	549616	1684.H17.gz43_263256
M00055806D:H03	ES 176	450252	1684.H23.gz43_263352
M00055806D:H06	ES 176	640520	1684.H24.gz43_263368
M00055807A:B10	ES 176	32812	1684.I03.gz43_263033
M00055807B:F05	ES 176	520320	1684.I12.gz43_263177
M00055807B:G10	ES 176	. 446789	1684,I14.gz43_263209
M00055807C:F05	ES 176	639441	1684,I17.gz43_263257
M00055807D:C04	ES 176	470602	1684.I21.gz43_263321
M00055808A:C06	ES 176	450741	1684.J05.gz43_263066
M00055808B;A04	ES 176	650782	1684.J08.gz43_263114
M00055808B:H07	ES 176	210839	1684.J14.gz43_263210
M00055808D:C12	ES 176	460517	1684.J20.gz43_263306

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CloneID	ES No	ClusterID	SequenceName
M00055808D:F09	ES 176	520320	1684.J24.gz43_263370
M00055809A:B10	ES 176	550370	1684.K05.gz43_263067
M00055809C:D01	ES 176	644901	1684.K17.gz43_263259
M00055809C:E11	ES 176	454772	1684.K19,gz43_263291
M00055809C:H10	ES 176	640479	1684.K22.gz43_263339
M00055810A;H03	ES 176	413277	1684.L06.gz43_263084
M00055810D:G04	ES 176	649995	1684.L19.gz43_263292
M00055811B:A03	ES 176	455665	1684.L23.gz43_263356
M00055811C:A01	ES 176	517327	1684.M01.gz43_263005
M00055811C:D12	ES 176	645015	1684.M03.gz43_263037
M00055811D:C02	ES 176	650231	1684.M07.gz43_263101
M00055811D:C12	ES 176	554742	1684.M09.gz43_263133
M00055811D:E01	ES 176	452828	1684.M10.gz43_263149
M00055812A:E01	ES 176	503491	1684.M15.gz43_263229
M00055812B;F03	ES 176	466016	1684.M19.gz43_263293
M00055812C:B07	ES 176	648752	1684.M24.gz43_263373
M00055812D:E03	ES 176	550604	1684.N09.gz43_263134
M00055813A:D10	ES 176	644147	1684.N12.gz43_263182
M00055813C:E03	ES 176	551683	1684.N17.gz43_263262
M00055813D;B12	ES 176	494271	1684,N19.gz43_263294
M00055814A:F02	ES 176	528134	1684.O01.gz43_263007
M00055814C:C07	ES 176	647336	1684.O11.gz43_263167
M00055815A:H12	ES 176	642850	1684,O24,gz43_263375
M00055815C:B03	ES 176	642099	1684.P04.gz43_263056
M00055815C:G05	ES 176	648221	1684.P08.gz43_263120
M00055815D:A11	ES 176	642962	1684.P09.gz43_263136
M00055815D:B02	ES 176	447015	1684.P10.gz43_263152
M00055815D:B03	ES 176	549640	1684.P11.gz43_263168
M00055816A:C04	ES 176	649035	1684.P17.gz43_263264
M00055816B:B07	ES 176	564854	1684.P23.gz43_263360
M00055816D:A10	ES 176	363172	1693.A07.gz43_213787
M00055816D:B11	ES 176	492242	1693.A09.gz43_213819
M00055816D:E10	ES 176	452863	1693,A13.gz43_213883
M00055817A;E05	ES 176	643397	1693.A17.gz43_213947
M00055817A;H07	ES 176	648580	1693,A19.gz43_213979
M00055817B:C04	ES 176	449542	1693,A23,gz43_214043
M00055818B:H03	ES 176	638837	1693,C10.gz43_213837
M00055819A:A06	ES 176	475624	1693.C23.gz43_214045
M00055819A:B10	ES 176	559854	1693.D01.gz43_213694
M00055822C:D09	ES 176	451966	1693.F18.gz43_213968
M00055823B:F02	ES 176	674526	1693.G08.gz43_213809
M00055825B:C11	ES 176	645155	1693.H03.gz43_213730
M00055825B:E03	ES 176	650204	1693.H04.gz43_213746
M00055825B:F09	ES 176	643054	1693,H06.gz43_213778

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Table 15			
CloneID	ES No	ClusterID	SequenceName
M00055825D:A03	ES 176	642962	1693.H13.gz43_213890
M00055825D:D11	ES 176	551681	1693.H20.gz43_214002
M00055826A:G04	ES 176	640464	1693.I06.gz43_213779
M00055826B:G07	ES 176	645746	1693.I13.gz43_213891
M00055826C:G06	ES 176	650487	1693.I19.gz43_213987
M00055827A:A12	ES 176	650579	1693.J01.gz43_213700
M00055827D:A01	ES 176	455716	1693.J19.gz43_213988
M00055827D:C02	ES 176	556286	1693.J22.gz43_214036
M00055827D:G11	ES 176	583915	1693.K03.gz43_213733
M00055828A:H10	ES 176	647425	1693,K07.gz43_213797
M00055828B:E10	ES 176	517280	1693.K10.gz43_213845
M00055829C:A07	ES 176	648567	1693.L02.gz43_213718
M00055829D:H10	ES 176	645146	1693.L15.gz43_213926
M00055830A:G10	ES 176	645073	1693,L17.gz43_213958
M00055830C:H10	ES 176	451134	1693.M06.gz43_213783
M00055831A:C06	ES 176	644781	1693.M15.gz43_213927
M00055831B:C04	ES 177	517237	1693.M23.gz43_214055
M00055832A:A08	ES 177	481864	1693.N08.gz43_213816
M00055832C:H09	ES 177	648328	1693.N19.gz43_213992
M00055833D:F11	ES 177	650912	1693.O14.gz43_213913
M00055834B:C11	ES 177	557741	1693.O20.gz43_214009
M00055835C:F08	ES 177	642054	1693.P21.gz43_214026
M00055836A:B12 ·	ES 177	645149	1694.A05.gz43_214139
M00055836C;D01	ES 177	452735	1694.A10.gz43_214219
M00055837A:B08	ES 177	447597	1694.A15.gz43_214299
M00055837A:D09	ES 177	644210	1694.A16.gz43_214315
M00055837A;F02	ES 177	489249	1694.A18.gz43_214347
M00055837A:H08	ES 177	645289	1694.A19.gz43_214363
M00055837B;E07	ES 177	644987	1694.A23.gz43_214427
M00055837D;D08	ES 177	642253	1694.B09.gz43_214204
M00055837D;G10	ES 177	504944	1694.B11.gz43_214236
M00055838A:A03	ES 177	643800	1694.B12.gz43_214252
M00055838A:B02	ES 177	448450	1694.B13.gz43_214268
M00055838B:D06	ES 177	467901	1694.B15.gz43 214300
M00055838B:G12	ES 177	644417	1694.B16.gz43 214316
M00055838B:H04	ES 177	644494	1694.B17.gz43 214332
M00055838C:A08	ES 177	447002	1694.B18.gz43_214348
M00055838C:B08	ES 177	643948	1694,B19.gz43_214364
M00055839B:A10	ES 177	643804	1694.C13.gz43_214269
M00055839B:C07	ES 177	648039	1694.C15.gz43_214301
M00055839B:E07	ES 177	558890	1694,C18.gz43_214349
M00055839B:H09	ES 177	537586	1694,C19.gz43_214365
M00055839C:B11	ES 177	467521	1694.C22.gz43 214413

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055839C:D06	ES 177	645948	1694.C23.gz43_214429
M00055840A:G06	ES 177	553145	1694.D08.gz43_214190
M00055840B:B02	ES 177	522869	1694.D11.gz43_214238
M00055840C:D06	ES 177	640618	1694.D17.gz43_214334
M00055840C:H06	ES 177	644548	1694.D18.gz43_214350
M00055840D:B03	ES 177	643977	1694.D20.gz43_214382
M00055841B:F09	ES 177	485237	1694.E09.gz43_214207
M00055841B:H03	ES 177	503546	1694.E10.gz43_214223
M00055841C:A03	ES 177	643843	1694.E12.gz43_214255
M00055841C:D05	ES 177	607430	1694.E15.gz43_214303
M00055841C:D11	ES 177	599759	1694.E17.gz43_214335
M00055841C:H04	ES 177	140648	1694.E20.gz43_214383
M00055841D:C11	ES 177	650528	1694.E21.gz43_214399
M00055842B:A04	ES 177	642079	1694.F05.gz43_214144
M00055842B:D04	ES 177	644063	1694.F09.gz43_214208
M00055842C:A11	ES 177	644548	1694.F11.gz43_214240
M00055842C:C03	ES 177	506901	1694.F12.gz43_214256
M00055842D:C02	ES 177	645505	1694.F15.gz43_214304
M00055842D:D07	ES 177	643809	1694.F17.gz43_214336
M00055842D:F07	ES 177	462557	1694.F19.gz43_214368
M00055843B:D10	ES 177	644212	1694.G04.gz43_214129
M00055843D:H01	ES 177	472119	1694.G15.gz43_214305
M00055844A:D03	ES 177	643825	1694.G18.gz43_214353
M00055844A:D07	ES 177	647577	1694.G19.gz43_214369
M00055844A:F11	ES 177	649068	1694.G20.gz43_214385
M00055844B:C12	ES 177	644075	1694.G24.gz43_214449
M00055844C:F01	ES 177	645290	1694.H06.gz43_214162
M00055844D:E11	ES 177	456728	1694.H10.gz43_214226
M00055844D:E12	ES 177	644314	1694.H11.gz43_214242
M00055844D:H09	ES 177	466697	1694.H16.gz43_214322
M00055845A:C03	ES 177	643999	1694.H19.gz43_214370
M00055845A:H10	ES 177	644569	1694.H21.gz43_214402
M00055845C:A11	ES 177	647448	1694.I05.gz43_214147
M00055845C:C12	ES 177	468222	1694.I08.gz43 214195
M00055845C:E02	ES 177	646420	1694.I10.gz43 214227
M00055845C:E05	ES 177	525456	1694.I11.gz43_214243
M00055845D:G11	ES 177	650276	1694.I16.gz43 214323
M00055846A:D07	ES 177	181364	1694.I18.gz43 214355
M00055846B:B12	ES 177	419751	1694.I23.gz43_214435
M00055846B:C12	ES 177	542282	1694.I24.gz43 214451
M00055846B:F11	ES 177	647191	1694.J01.gz43 214084
M00055846D:G08	ES 177	647383	1694.J12.gz43_214260
M00055846D:G09	ES 177	645681	1694.J13.gz43 214276
M00055846D:G11	ES 177	640171	1694.J14.gz43 214292

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Table 15			
CloneID	ES No	ClusterID	SequenceName
M00055847B:G12	ES 177	651131	1694,J19.gz43_214372
M00055847C:A11	ES 177	646459	1694.J21.gz43_214404
M00055847C:C01	ES 177	646810	1694.J22.gz43_214420
M00055847C:H09	ES 177	452976	1694.J24.gz43_214452
M00055848A:E07	ES 177	645681	1694.K06.gz43_214165
M00055848B:C03	ES 177	647383	1694.K13.gz43_214277
M00055848C:A02	ES 177	645197	1694.K15.gz43_214309
M00055848C:G07	ES 177	558981	1694.K17.gz43_214341
M00055848C:H06	ES 177	646248	1694,K19.gz43_214373
M00055849C:G07	ES 177	547841	1694.L06.gz43_214166
M00055849D:B04	ES 177	450949	1694.L08.gz43_214198
M00055849D:H09	ES 177	469511	1694,L13.gz43_214278
M00055850C:D01	ES 177	608873	1694,L21.gz43_214406
M00055850C:G05	ES 177	647318	1694.L22.gz43_214422
M00055851A:C03	ES 177	466265	1694.M06.gz43_214167
M00055851A:C09	ES 177	649842	1694,M07.gz43_214183
M00055851A:G11	ES 177	97507	1694.M12.gz43_214263
M00055851A:H10	ES 177	647522	1694,M14.gz43_214295
M00055851B:B09	ES 177	562229	1694,M16.gz43_214327
M00055851B:G02	ES 177	639295	1694.M18.gz43_214359
M00055851B:G10	ES 177	425923	1694,M19.gz43_214375
M00055851C:F12	ES 177	643594	1694.M23.gz43_214439
M00055851C:H05	ES 177	472801	1694.M24.gz43_214455
M00055852A:C12	ES 177	639341	1694.N09.gz43_214216
M00055852B;F10	ES 177	562813	1694.N12.gz43_214264
M00055852B:G09	ES 177	642293	1694.N14.gz43 214296
M00055852B:H04	ES 177	457661	1694.N16.gz43 214328
M00055852D:B11	ES 177	644708	1694.N20.gz43_214392
M00055852D:G12	ES 177	649611	1694.N21.gz43_214408
M00055853B:H06	ES 177	647578	1694.O04.gz43_214137
M00055853C:C12	ES 177	603388	1694.O08.gz43_214201
M00055853C:H03	ES 177	647577	1694,O10.gz43_214233
M00055853D:A07	ES 177	650492	1694.O11.gz43_214249
M00055853D:B04	ES 177	643130	1694.O13.gz43_214281
M00055853D:C07	ES 177	650217	1694.O15.gz43 214313
M00055854A:B07	ES 177	592122	1694.O17.gz43_214345
M00055854A;D01	ES 177	639510	1694.O19.gz43_214377
M00055854A:E03	ES 177	645145	1694.O21.gz43 214409
M00055854A:E04	ES 177	638854	1694.O22.gz43_214425
M00055854C:C07	ES 177	549624	1694,P04,gz43 214138
M00055854C:E03	ES 177	570939	1694.P05.gz43 214154
M00055854C:H11	ES 177	471364	1694.P09.gz43 214218
M00055855A:B11	ES 177	404816	1694.P14.gz43 214298
M00055855A;G05	ES 177	462659	1694.P16.gz43 214330

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Table 15			
CloneID	ES No	ClusterID	SequenceName
M00055855B;B11	ES 177	643909	1694.P20.gz43_214394
M00055855B:D12	ES 177	465576	1694.P21.gz43_214410
M00055855C:F11	ES 177	603388	1694,P23.gz43_214442
M00055855D:D12	ES 177	509027	1695.A04.gz43_214911
M00055855D:G08	ES 177	644442	1695.A07.gz43_214959
M00055856A:C06	ES 177	564854	1695.A08.gz43_214975
M00055856A:D12	ES 177	644149	1695.A09.gz43_214991
M00055856C:F07	ES 177	550267	1695.A20.gz43_215167
M00055857C:D09	ES 177	646372	1695.B08.gz43_214976
M00055858A:G07	ES 177	642070	1695.B24.gz43_215232
M00055860D:E04	ES 177	596882	1695.D05.gz43_214930
M00055861B:F04	ES 177	645848	1695.D08.gz43_214978
M00055861C:G03	ES 177	645538	1695.D13.gz43_215058
M00055862A:C01	ES 177	140224	1695.D21.gz43_215186
M00055862D:B02	ES 177	640147	1695.E09.gz43_214995
M00055862D:D06	ES 177	449500	1695,E11,gz43_215027
M00055863B:C07	ES 177	644030	1695.E20.gz43_215171
M00055863D:D09	ES 177	644047	1695.F07.gz43_214964
M00055864A:C09	ES 177	644047	1695.F13.gz43_215060
M00055864A:E11	ES 177	650773	1695.F14.gz43_215076
M00055864B:C09	ES 177	561877	1695.F18.gz43_215140
M00055865C:G11	ES 177	649660	1695.G23.gz43_215221
M00055865C:H06	ES 177	650517	1695,G24,gz43_215237
M00055866A:G10	ES 177	397399	1695.H07.gz43_214966
M00055866C:G09	ES 177	644435	1695.H17.gz43_215126
M00055866C:H06	ES 177	644410	1695.H18.gz43_215142
M00055867A:B02	ES 177	645288	1695.H23.gz43_215222
M00055868B;B04	ES 177	649436	1695.I16.gz43_215111
M00055868C:F02	ES 177	562059	1695.I24.gz43_215239
M00055868D:D08	ES 177	469437	1695.J05.gz43_214936
M00055869C:G06	ES 177	449936	1695,J14.gz43_215080
M00055869D:A07	ES 177	642411	1695.J16.gz43_215112
M00055870B:D04	ES 177	649846	1695.J24.gz43_215240
M00055871A:H06	ES 177	524261	1695.K09.gz43 215001
M00055871B:B03	ES 177	648819	1695.K11.gz43_215033
M00055871C:C07	ES 177	643099	1695.K21.gz43 215193
M00055871C:C10	ES 177	644801	1695.K22.gz43 215209
M00055872A:C08	ES 177	644755	1695.L11.gz43_215034
M00055872A:D08	ES 177	644830	1695.L12.gz43_215050
M00055872A:E05	ES 177	505858	1695.L13.gz43_215066
M00055872A:E11	ES 177	640534	1695.L15.gz43_215098
M00055872D:D12	ES 177	447035	1695.M08.gz43_214987
M00055873A:E03	ES 177	643451	1695.M16.gz43_215115
M00055873D:E04	ES 177	643488	1695.N10.gz43 215020

Table 13

CloneID	ES No	ClusterID	SequenceName
M00055874A:F06	ES 177	645497	1695.N18.gz43_215148
M00055875A;G05	ES 177	554646	1695.O14.gz43_215085
M00055875B;E09	ES 177	646633	1695.P03.gz43_214910
M00055875C:C07	ES 177	516512	1695.P11.gz43_215038
M00055875D:A07	ES 177	. 640136	1695.P20.gz43_215182
M00055875D:D01	ES 177	643333	1695.P22.gz43_215214
M00055875D:H10	ES 177	650179	1696.A01.gz43_215247
M00055877A:H04	ES 177	645638	1696.A07.gz43_215343
M00055877B:B05	ES 177	484991	1696.A09.gz43_215375
M00055877D:C05	ES 177	643233	1696,A21.gz43_215567
M00055878C:C02	ES 177	593715	1696.B16.gz43_215488
M00055879A:D04	ES 177	643934	1696.C11.gz43_215409
M00055879A:G03	ES 177	450001	1696.C12.gz43_215425
M00055880B:A04	ES 177	642911	1696.D03.gz43_215282
M00055880B:A06	ES 177	447807	1696.D04.gz43_215298
M00055880C:F07	ES 177	645215	1696.D15.gz43_215474
M00055881A:A08	ES 177	447221	1696.E05.gz43_215315
M00055881A:E10	ES 177	479868	1696.E12.gz43_215427
M00055881A;G07	ES 177	446225	1696.E13.gz43_215443
M00055881D:A02	ES 177	648063	1696.E22.gz43_215587
M00055882C:A06	ES 177	450559	1696.F14.gz43_215460
M00055882D:B02	ES 177	640400	1696.F21.gz43_215572
M00055883A:C02	ES 177	179760	1696.G05.gz43_215317
M00055883A:C10	ES 177	459274	1696.G06.gz43_215333
M00055883A:H01	ES 177	643745	1696.G11.gz43_215413
M00055883D:B06	ES 177	644468	1696.G21.gz43_215573
M00055884B:F10	ES 177	451670	1696.H17.gz43_215510
M00055884B:H07	ES 177	552628	1696.H18.gz43_215526
M00055884C:B07	ES 177	643046	1696.H21.gz43_215574
M00055884D:F07	ES 177	345761	1696.I06.gz43_215335
M00055885A:D05	ES 177	558222	1696.I11.gz43_215415
M00055885B:A11	ES 177	555172	1696.I15.gz43_215479
M00055885B:B04	ES 178	586992	1696.I16.gz43_215495
M00055885C:B07	ES 178	552201	1696.J01.gz43_215256
M00055886A:B06	ES 178	643126	1696.J12.gz43_215432
M00055886D:E11	ES 178	557792	1696.K11.gz43_215417
M00055887A:F07	ES 178	644099	1696.K18.gz43_215529
M00055887D:C11	ES 178	643248	1696.L14.gz43_215466
M00055888C:F07	ES 178	639256	1696.M04.gz43_215307
M00055888C:G09	ES 178	640356	1696.M06.gz43_215339
M00055889B:E12	ES 178	570248	1696.M23.gz43_215611
M00055890A:A06	ES 178	464171	1696.N11.gz43_215420
M00055890A:D01	ES 178	549114	1696.N14.gz43_215468

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055890A:G12	ES 178	649068	1696.N21.gz43_215580
M00055890A:H11	ES 178	643724	1696.N23.gz43_215612
M00055891B:F09	ES 178	643513	1696.P02.gz43_215278
M00055892A:F04	ES 178	649027	1696.P23.gz43 215614
M00055892B:D02	ES 178	643366	1696.P24.gz43 215630
M00055893B:C05	ES 178	650476	1705.A11.gz43 215791
M00055894A:H08	ES 178	374340	1705.A22.gz43 215967
M00055894B:E09	ES 178	394373	1705.B03.gz43 215664
M00055896A:G01	ES 178	648576	1705.B11.gz43 215792
M00055896B:C06	ES 178	643948	1705.B14.gz43 215840
M00055896B;H04	ES 178	640762	1705.B18.gz43 215904
M00055896C:H10	ES 178	553951	1705.B23.gz43_215984
M00055896D:G10	ES 178	557361	1705.C03.gz43_215665
M00055898B:E07	ES 178	485237	1705.C11.gz43 215793
M00055899A:B03	ES 178	645844	1705.C18.gz43_215905
M00055899C:H01	ES 178	474577	1705.D05.gz43_215698
M00055900A:F08	ES 178	644390	1705.D16.gz43_215874
M00055900B;E12	ES 178	446728	1705.D20.gz43_215938
M00055900D:D03	ES 178	526575	1705.E01.gz43_215635
M00055901B:A02	ES 178	647522	1705.E10.gz43_215779
M00055901C:A01	ES 178	553720	1705.E16.gz43_215875
M00055901C:C10	ES 178	641087	1705.E18.gz43_215907
M00055901C:D04	ES 178	642940	1705.E19.gz43_215923
M00055902A:H11	ES 178	643513	1705.F06.gz43_215716
M00055902D:A07	ES 178	557947	1705.F14.gz43_215844
M00055902D:H10	ES 178	482163	1705.F18.gz43_215908
M00055903B:C10	ES 178	644063	1705.F24.gz43_216004
M00055903B:E05	ES 178	554166	1705,G03.gz43_215669
M00055904B:B01	ES 178	643914	1705.G12.gz43_215813
M00055904C:A07	ES 178	643809	1705,G15.gz43_215861
M00055904D:A06	ES 178	643808	1705.G18.gz43_215909
M00055905A:F11	ES 178	644354	1705.G24.gz43_216005
M00055905B:H01	ES 178	639194	1705,H06.gz43_215718
M00055905D:B06	ES 178	605596	1705,H13.gz43_215830
M00055905D:H02	ES 178	550242	1705.H17.gz43_215894
M00055906C:F03	ES 178	641562	1705.H23.gz43_215990
M00055906D:G03	ES 178	454720	1705.I06.gz43_215719
M00055907D:D06	ES 178	449994	1705.J06.gz43_215720
M00055908D:F09	ES 178	644364	1705.K04.gz43_215689
M00055909B:B12	ES 178	638869	1705,K10.gz43_215785
M00055909B:G10	ES 178	643089	1705,K16.gz43_215881
M00055909C:E08	ES 178	447326	1705.K20.gz43_215945
M00055909C:G09	ES 178	645781	1705.K21.gz43_215961
M00055909D:A09	ES 178	638919	1705.K22.gz43 215977

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055909D:E01	ES 178	89082	1705.K24.gz43_216009
M00055910B:B11	ES 178	642985	1705.L11.gz43_215802
M00055910B:H12	ES 178	639240	1705.L16.gz43_215882
M00055910C:G01	ES 178	468147	1705.L19.gz43_215930
M00055910C:G04	ES 178	685001	1705.L20.gz43_215946
M00055910D:A03	ES 178	643825	1705.L21.gz43_215962
M00055911B:E06	ES 178	644242	1705.M05.gz43_215707
M00055912A:F08	ES 178	557675	1705.M16.gz43_215883
M00055912B:E05	ES 178	648237	1705.M20.gz43_215947
M00055912C:C08	ES 178	640756	1705.M22.gz43_215979
M00055912D:F04	ES 178	644407	1705.N05.gz43_215708
M00055913A:G07	ES 178	649312	1705.N12.gz43_215820
M00055913B:D05	ES 178	644190	1705.N17.gz43_215900
M00055914A:A03	ES 178	449737	1705.O02.gz43_215661
M00055914C:D12	ES 178	643300	1705.O09.gz43_215773
M00055914C:G01	ES 178	644108	1705.O11.gz43_215805
M00055914C:G11	ES 178	641645	1705.O12.gz43_215821
M00055914D:A08	ES 178	564134	1705.O13.gz43_215837
M00055914D:G06	ES 178	553587	1705.O16.gz43_215885
M00055915A:A06	ES 178	639413	1705.O17.gz43_215901
M00055915B:D08	ES 178	447212	1705.O21.gz43_215965
M00055915C:B09	ES 178	643991	1705.O24.gz43_216013
M00055915D:A07	ES 178	640178	1705.P05.gz43_215710
M00055916A:E12	ES 178	644236	1705.P12.gz43_215822
M00055916B:C02	ES 178	649170	1705.P15.gz43_215870
M00055917B:D02	ES 178	642911	1706.A08.gz43_216139
M00055917C:F04	ES 178	549994	1706.A13.gz43_216219
M00055917D:E07	ES 178	644301	1706.A16.gz43_216267
M00055918A:F10	ES 178	645028	1706.A20.gz43_216331
M00055918B:B07	ES 178	643936	1706.A24.gz43_216395
M00055918B:B10	ES 178	459881	1706.B01.gz43_216028
M00055918B:F10	ES 178	557175	1706.B03.gz43_216060
M00055918C:C04	ES 178	651131	1706.B05.gz43_216092
M00055918C:E02	ES 178	642166	1706.B07.gz43_216124
M00055919A:A06	ES 178	647952	1706.B12.gz43_216204
M00055919B:H11	ES 178	562021	1706.B19.gz43 216316
M00055919D:H07	ES 178	644554	1706.C01.gz43 216029
M00055920A:H10	ES 178	640977	1706.C05.gz43 216093
M00055920B;F02	ES 178	644410	1706.C09.gz43 216157
M00055921A:E06	ES 178	188079	1706.C21.gz43_216349
M00055922A:F05	ES 178	465576	1706.D13.gz43 216222
M00055922A:G07	ES 178	650900	1706.D15.gz43 216254
M00055922B;A06	ES 178	447426	1706.D17.gz43_216286
M00055922B:G09	ES 178	649085	1706.D21.gz43 216350

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Table 15			
CloneID	ES No	ClusterID	SequenceName
M00055922D:G10	ES 178	419009	1706.E06.gz43_216111
M00055923B:C07	ES 178	643170	1706,E10.gz43_216175
M00055924B:D02	ES 178	643627	1706.E22.gz43_216367
M00055924B:E12	ES 178	464129	1706.F01.gz43_216032
M00055924C:A09	ES 178	641484	1706,F03.gz43 216064
M00055925A:C04	ES 178	450840	1706.F11.gz43_216192
M00055925B:D09	ES 178	639154	1706.F20.gz43_216336
M00055925B:D10	ES 178	643332	1706.F21.gz43_216352
M00055926B;A05	ES 178	447123	1706.G12.gz43_216209
M00055926C:A11	ES 178	207552	1706.G18.gz43_216305
M00055927B:D06	ES 178	642308	1706.H05.gz43_216098
M00055927D:E11	ES 178	643239	1706.H10.gz43_216178
M00055928A:C06	ES 178	639038	1706.H15.gz43_216258
M00055928A:E11	ES 178	610893	1706.H17.gz43_216290
M00055928B:E11	ES 178	643383	1706.H21.gz43_216354
M00055929A:C11	ES 178	649954	1706.I14.gz43_216243
M00055929B:E01	ES 178	462089	1706.I17.gz43_216291
M00055929D:F09	ES 178	649954	1706.I24.gz43_216403
M00055930A:G04	ES 178	644629	1706.J05.gz43_216100
M00055930C:D03	ES 178	650938	1706.J09.gz43_216164
M00055930C:F06	ES 178	639596	1706.J10.gz43_216180
M00055930C:H05	ES 178	556925	1706.J15.gz43_216260
M00055930D:B05	ES 178	447520	1706.J16.gz43_216276
M00055930D:B09	ES 178	613626	1706.J17.gz43_216292
M00055930D:F05	ES 178	513211	1706.J21.gz43 216356
M00055933A:E05	ES 178	638808	1706.L07.gz43 216134
M00055933B:B11	ES 178	605114	1706.L10.gz43 216182
M00055933B:G01	ES 178	650245	1706.L11.gz43_216198
M00055934C:E06	ES 178	446795	1706.M06.gz43_216119
M00055934D:H09	ES 178	650470	1706.M12.gz43_216215
M00055935B:H06	ES 178	479604	1706.M20.gz43_216343
M00055935C:E03	ES 178	640672	1706.M23.gz43_216391
M00055936C:B05	ES 178	646596	1706.N07.gz43_216136
M00055936C:D03	ES 178	648379	1706.N08.gz43_216152
M00055936D:F03	ES 178	456920	1706.N11.gz43_216200
M00055937A:F07	ES 178	645092	1706.N17.gz43_216296
M00055937C:C08	ES 178	641716	1706.N21.gz43_216360
M00055939A;B12	ES 178	644732	1706.P02.gz43 216058
M00055939B;A11	ES 178	639901	1706.P07.gz43 216138
M00055939D:D07	ES 178	559905	1706.P13.gz43_216234
M00055940B:A08	ES 178	649873	1706.P19.gz43_216330
M00055941B:A04	ES 178	524546	1707.A12.gz43_216604
M00055942B:H10	ES 178	646317	1707.B04.gz43_216477
M00055942D:A01	ES 178	648782	1707.B12.gz43_216605

Table 13

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	CloneID	ES No	ClusterID	SequenceName
	M00055942D:E05	ES 178	645809	1707.B16.gz43_216669
1	M00055943B:E01	ES 178	447805	1707.B24.gz43_216797
1	M00055945C:C02	ES 178	648588	1707.D13.gz43_216623
Ī	M00055945C:D09	ES 178	648555	1707.D14.gz43 216639
1	M00055945D:D01	ES 178	461363	1707.D17.gz43 216687
Ī	M00055945D:E08	ES 178	644720	1707.D18.gz43 216703
1	M00055947C:E09	ES 178	393599	1707.E20.gz43_216736
1	M00055949A:E09	ES 178	499693	1707.F02.gz43 216449
1	M00055949D:B07	ES 178	649549	1707.F06.gz43_216513
1	M00055951B:A07	ES 178	644983	1707.G03.gz43 216466
Ī	M00055952C:E08	ES 178	650935	1707.G21.gz43_216754
1	M00055953D:B08	ES 178	642105	1707.H10.gz43_216579
1	M00055954C:G07	ES 178	531529	1707.H19.gz43_216723
1	M00055954D:B04	ES 178	561777	1707.H22.gz43 216771
1	M00055955B:C06	ES 178	460245	1707.I08.gz43 216548
1	M00055955C:H05	ES 178	650370	1707.I14.gz43_216644
Ī	M00055956D:B01	ES 178	550237	1707.J02.gz43_216453
Ī	M00055956D:C11	ES 178	447224	1707.J04.gz43_216485
1	M00055957B:F11	ES 178	452316	1707.J14.gz43_216645
Ī	M00055957C:F05	ES 178	644568	1707.J18.gz43_216709
[]	M00055958C:E03	ES 178	643968	1707.K05.gz43_216502
	M00055958D:F02	ES 178	647185	1707.K10.gz43_216582
Ī	M00055961D:F06	ES 178	647312	1707.L18.gz43_216711
Ī	M00055963B:C02	ES 178	555210	1707.L22.gz43_216775
1	M00055965A:G06	ES 178	642653	1707.M24.gz43_216808
	M00055965D:G01	ES 178	472101	1707.N12.gz43_216617
Ī	M00055966B:H09	ES 178	646190	1707.N18.gz43_216713
Ī	M00055966C:D03	ES 178	645508	1707.N20.gz43_216745
Ī	M00055967A:F11	ES 178	513619	1707.O07.gz43_216538
[	M00055967A:G01	ES 178	646061	1707.O08.gz43_216554
1	M00055967B:B12	ES 178	645273	1707.O09.gz43_216570
[	M00055967B:F07	ES 178	645900	1707.O13.gz43_216634
	M00055967D:C05	ES 178	383609	1707.O23.gz43_216794
1	M00055968D:E03	ES 178	643162	1707.P18.gz43_216715
	M00055969A:F02	ES 178	694643	1707.P24.gz43_216811
	M00055969B:B05	ES 178	640672	1708.A04.gz43_216860
	M00055969D:B08	ES 178	645289	1708.A11.gz43_216972
	M00055969D:F08	ES 178	642955	1708.A12.gz43_216988
	M00055970C:A05	ES 178	647058	1708.A21.gz43_217132
	M00055970D:F09	ES 178	645894	1708.B10.gz43_216957
	M00055971B:A11	ES 178	649082	1708.B14.gz43_217021
	M00055972C:C09	ES 178	557852	1708.C06.gz43_216894
	M00055972C:F02	ES 178	558052	1708.C07.gz43_216910
1	M00055973C:F10	ES 178	647211	1708.C24.gz43_217182
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CloneID	ES No	ClusterID	SequenceName
M00055973D:H11	ES 178	452762	1708.D08.gz43_216927
M00055974D:B01	ES 178	645264	1708.D18.gz43_217087
M00055976A:D04	ES 178	645926	1708.E22.gz43_217152
M00055976B:F01	ES 178	647280	1708.F05.gz43_216881
M00055977A:G09	ES 178	647333	1708.F19.gz43_217105
M00055977B:F06	ES 179	643723	1708.F21.gz43_217137
M00055978A:H03	ES 179	642058	1708.G11.gz43_216978
M00055978B:F01	ES 179	484355	1708.G16.gz43_217058
M00055979B:G07	ES 179	640092	1708.H12.gz43_216995
M00055979C:B07	ES 179	640204	1708.H15.gz43_217043
M00055980A:H06	ES 179	647539	1708.H23.gz43_217171
M00055980B:B07	ES 179	647924	1708.I01.gz43_216820
M00055980C:G12	ES 179	644354	1708.I08.gz43_216932
M00055981D:B07	ES 179	650513	1708.J07.gz43_216917
M00055982A:G06	ES 179	646013	1708.J17.gz43_217077
M00055982C:A12	ES 179	650018	1708.J24.gz43_217189
M00055982C:H01	ES 179	644226	1708.K04.gz43_216870
M00055983C:C09	ES 179	451709	1708.K14.gz43_217030
M00055983D:A09	ES 179	477790	1708.K18.gz43_217094
M00055984A:F05	ES 179	645848	1708.K24.gz43_217190
M00055984D:E04	ES 179	645746	1708.L08.gz43_216935
M00055985A:B06	ES 179	645288	1708.L10.gz43_216967
M00055985B:C02	ES 179	645427	1708.L13.gz43_217015
M00055985B:G12	ES 179	646126	1708.L16.gz43_217063
M00055985D:E09	ES 179	641484	1708.M02.gz43_216840
M00055986A:F05	ES 179	647539	1708.M09.gz43_216952
M00055987B:F07	ES 179	465209	1708.N06.gz43_216905
M00055988A:A12	ES 179	640179	1708.N15.gz43_217049
M00055988A:E05	ES 179	639359	1708.N21.gz43_217145
M00055989A:C09	ES 179	560349	1708.O11.gz43_216986
M00055990A:F07	ES 179	645920	1708.P01.gz43_216827
M00055990D:B02	ES 179	558452	1708.P12.gz43_217003
M00055990D:G09	ES 179	582549	1708.P15.gz43_217051
M00055991A:D10	ES 179	649309	1708.P18.gz43_217099
M00055991A:H09	ES 179	450123	1708.P20.gz43_217131
M00055991C:H11	ES 179	646212	1708.P24.gz43_217195
M00055992A:D08	ES 179	645662	1717.A08.gz43_218513
M00055993A:E02	ES 179	643288	1717.A19.gz43_218689
M00055993D:F12	ES 179	649556	1717.B06.gz43_218482
M00055994A:G04	ES 179	603388	1717.B11.gz43_218562
M00055994B:B12	ES 179	651119	1717.B15.gz43_218626
M00055994C:E06	ES 179	642887	1717.B23.gz43_218754
M00055994C:F05	ES 179	639377	1717.B24.gz43_218770

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055995A:G09	ES 179	643808	1717.C14.gz43_218611
M00055995B:A02	ES 179	477521	1717.C15.gz43_218627
M00055995C:F05	ES 179	650534	1717.C20.gz43_218707
M00055996A:C10	ES 179	466277	1717.D02.gz43_218420
M00055996B:H04	ES 179	466271	1717.D08.gz43_218516
M00055996C:A02	ES 179	650018	1717.D09.gz43_218532
M00055996C:A08	ES 179	640645	1717.D10.gz43_218548
M00055996D:D07	ES 179	558856	1717.D13.gz43_218596
M00055996D:G02	ES 179	587106	1717.D15.gz43_218628
M00055997A:B08	ES 179	642000	1717.D19.gz43_218692
M00055997A:C08	ES 179	634365	1717.D20.gz43_218708
M00055997A:F04	ES 179	552846	1717.D23.gz43_218756
M00055997D:A11	ES 179	541214	1717.E11.gz43_218565
M00055998A:F06	ES 179	644990	1717.E20.gz43_218709
M00055998B:A04	ES 179	644612	1717.E22.gz43_218741
M00055998B:D06	ES 179	560860	1717.F01.gz43_218406
M00055998B:G03	ES 179	402100	1717.F03.gz43_218438
M00055998C:D04	ES 179	513888	1717.F05.gz43_218470
M00055998D:G05	ES 179	645076	1717.F18.gz43_218678
M00055999A:A04	ES 179	478025	1717.F19.gz43_218694
M00055999A:B07	ES 179	207530	1717.F20.gz43_218710
M00055999A:H10	ES 179	646387	1717.F22.gz43_218742
M00055999C:A01	ES 179	642881	1717.G04.gz43_218455
M00055999D:C05	ES 179	549889	1717.G12.gz43_218583
M00055999D:H04	ES 179	644210	1717.G17.gz43_218663
M00056000A:B02	ES 179	447822	1717.G19.gz43_218695
M00056000A:E11	ES 179	647443	1717.G22.gz43_218743
M00056000C:D09	ES 179	649259	1717.H08.gz43_218520
M00056001C:E09	ES 179	562221	1717.H20.gz43_218712
M00056001C:F07	ES 179	447556	1717.H21.gz43_218728
M00056001D:B06	ES 179	641736	1717.I03.gz43_218441
M00056001D:G12	ES 179	643971	1717.I05.gz43_218473
M00056002A:A03	ES 179	644611	1717.I06.gz43_218489
M00056002A:H06	ES 179	559400	1717.109.gz43_218537
M00056002D:E09	ES 179	419443	1717.I15.gz43_218633
M00056003A:A08	ES 179	449261	1717.I17.gz43_218665
M00056003A:B07	ES 179	641580	1717.I19.gz43_218697
M00056003A:C01	ES 179	645197	1717.I20.gz43_218713
M00056003B:E11	ES 179	618670	1717.J02.gz43_218426
M00056003B:G11	ES 179	464498	1717.J03.gz43_218442
M00056003C:C09	ES 179	644824	1717.J05.gz43_218474
M00056003C:H10	ES 179	645162	1717.J07.gz43_218506
M00056004A:E12	ES 179	451469	1717.J13.gz43_218602
M00056004B:F04	ES 179	643498	1717.J17.gz43_218666

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Table 13			<u>-</u>
CloneID	ES No	ClusterID	SequenceName
M00056004D:F03	ES 179	452775	1717.K03.gz43_218443
M00056004D:F12	ES 179	444454	1717.K04.gz43_218459
M00056005B:H08	ES 179	641167	1717.K20.gz43_218715
M00056005C:F11	ES 179	561836	1717.L03.gz43_218444
M00056005D:C04	ES 179	470462	1717.L07.gz43_218508
M00056005D:F12	ES 179	362177	1717.L10.gz43_218556
M00056006A:E04	ES 179	643440	1717.L15.gz43_218636
M00056006A;G09	ES 179	532307	1717.L18.gz43_218684
M00056006B;A10	ES 179	637977	1717.L19.gz43_218700
M00056006B;B05	ES 179	592941	1717.L20.gz43_218716
M00056006B:C12	ES 179	411885	1717.L24.gz43_218780
M00056006D:C06	ES 179	643239	1717.M06.gz43_218493
M00056006D:D05	ES 179	639940	1717.M07.gz43_218509
M00056006D:E03	ES 179	643477	1717.M10.gz43_218557
M00056006D:F05	ES 179	651083	1717.M11.gz43_218573
M00056007A:B02	ES 179	555736	1717.M16.gz43_218653
M00056007A;G07	ES 179	647194	1717.M22.gz43_218749
M00056007B:C10	ES 179	449713	1717.M24.gz43_218781
M00056007B:E08	ES 179	649701	1717.N03.gz43_218446
M00056007C:A03	ES 179	448418	1717.N05.gz43_218478
M00056007D:F07	ES 179	644022	1717.N13.gz43_218606
M00056007D:H12	ES 179	262760	1717.N14.gz43_218622
M00056008A;H03	ES 179	675319	1717.N17.gz43 218670
M00056008B:E04	ES 179	641283	1717.N21.gz43 218734
M00056008B:G01	ES 179	456627	1717.N24.gz43 218782
M00056008B:G05	ES 179	447802	1717.O02.gz43_218431
M00056008C:D04	ES 179	639178	1717.O04.gz43_218463
M00056008C:D08	ES 179	644030	1717.O05.gz43_218479
M00056008D:D02	ES 179	644612	1717.O13.gz43_218607
M00056008D:D03	ES 179	642260	1717.O14.gz43_218623
M00056009A:H08	ES 179	557895	1717.O18.gz43_218687
M00056009C:F09	ES 179	649860	1717.O23.gz43_218767
M00056009D:A02	ES 179	646143	1717.O24.gz43_218783
M00056009D:C12	ES 179	640709	1717.P01.gz43_218416
M00056010A;E05	ES 179	644972	1717.P05.gz43 218480
M00056011A:C11	ES 179	644949	1717.P13.gz43 218608
M00056011B:A06	ES 179	517274	1717.P17.gz43 218672
M00056011B:E10	ES 179	489426	1717.P19.gz43 218704
M00056011C:D04	ES 179	555172	1717.P22.gz43 218752
M00056013A:C09	ES 179	538808	1718.B12.gz43 218965
M00056013D:D07	ES 179	640181	1718.B21.gz43_219109
M00056016C:F11	ES 179	643744	1718.D04.gz43 218839
M00056016D:E02	ES 179	644967	1718.D07.gz43 218887
M00056017A:B08	ES 179	643975	1718,D09,gz43 218919

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CloneID	ES No	ClusterID	SequenceName
M00056019B;A01	ES 179	643510	1718.F02.gz43 218809
M00056020B:C09	ES 179	650348	1718,F23.gz43 219145
M00056020C:H03	ES 179	650391	1718.G06.gz43 218874
M00056020D:D03	ES 179	545509	1718.G08.gz43 218906
M00056021A;H06	ES 179	557344	1718.G17.gz43 219050
M00056021C:H08	ES 179	639629	1718.H03.gz43 218827
M00056023D:D08	ES 179	650749	1718.I21.gz43_219116
M00056026B:H01	ES 179	644982	1718.K18.gz43_219070
M00056028D:F06	ES 179	645883	1718.M11.gz43_218960
M00056029B:H03	ES 179	467057	1718.M20.gz43_219104
M00056029D:D08	ES 179	447659	1718.N06.gz43_218881
M00056031D:E06	ES 179	495942	1718.O14.gz43_219010
M00056032B:H08	ES 179	397399	1718.P04.gz43_218851
M00056033A:C11	ES 179	425455	1718.P13.gz43_218995
M00056033C:H09	ES 179	451993	1718.P24.gz43_219171
M00056033D:F06	ES 179	645973	1719.A03.gz43_219204
M00056033D:G07	ES 179	642198	1719.A04.gz43_219220
M00056034A:G10	ES 179	646048	1719.A10.gz43_219316
M00056034C:D07	ES 179	452325	1719.A17.gz43_219428
M00056034C:H09	ES 179	648390	1719.A21.gz43_219492
M00056034D:E09	ES 179	640559	1719.A23.gz43_219524
M00056035B:A04	ES 179	550376	1719.B03.gz43_219205
M00056035B:D11	ES 179	566745	1719.B06.gz43_219253
M00056035B:E10	ES 179	549786	1719.B07.gz43_219269
M00056035C:H11	ES 179	452504	1719.B15.gz43_219397
M00056035D:C08	ES 179	447645	1719.B17.gz43_219429
M00056035D:G10	ES 179	467597	1719.B21.gz43_219493
M00056036B:B01	ES 179	648045	1719.C01.gz43_219174
M00056036B:F02	ES 179	408428	1719.C05.gz43_219238
M00056036D:B06	ES 179	468689	1719.C14.gz43_219382
M00056036D:D01	ES 179	452729	1719.C15.gz43_219398
M00056036D:F05	ES 179	644054	1719.C17.gz43_219430
M00056037A:D11	ES 179	645530	1719.C20.gz43_219478
M00056037A;E10	ES 179	645707	1719.C22.gz43_219510
M00056037B:G02	ES 179	646088	1719.D03.gz43_219207
M00056037C:D06	ES 179	476077	1719.D09.gz43_219303
M00056037C:G11	ES 179	· 638912	1719.D10.gz43_219319
M00056038B:G08	ES 179	481614	1719.D15.gz43_219399
M00056038D:F03	ES 179	468330	1719.D21.gz43_219495
M00056039A:A02	ES 179	550365	1719,D23.gz43_219527
M00056039A:F04	ES 179	647976	1719.E01.gz43_219176
M00056039B:C03	ES 179	649293	1719,E04.gz43_219224
M00056039C:D05	ES 179	453804	1719,E08.gz43_219288
M00056039C:G05	ES 179	645973	1719.E11.gz43_219336

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Table 15			
CloneID	ES No	ClusterID	SequenceName
M00056039C;H01	ES 179	642852	1719.E12.gz43_219352
M00056040A;E09	ES 179	647144	1719.E15.gz43_219400
M00056040C:A03	ES 179	649118	1719.E20.gz43_219480
M00056040C:B03	ES 179	454438	1719.E21.gz43 219496
M00056040C:C12	ES 179	646687	1719.E22.gz43_219512
M00056041A:C07	ES 179	479722	1719.F05.gz43_219241
M00056041B;F03	ES 179	644817	1719.F10.gz43_219321
M00056042B:A03	ES 179	558432	1719.F23.gz43_219529
M00056042B:B05	ES 179	483266	1719.F24.gz43_219545
M00056043A:A09	ES 179	460727	1719.G13.gz43_219370
M00056043A:H12	ES 179	556325	1719.G16.gz43_219418
M00056043B:E03	ES 179	643897	1719.G19.gz43_219466
M00056043C:G03	ES 179	490898	1719.G24.gz43_219546
M00056043D:E03	ES 179	641193	1719.H04.gz43_219227
M00056044B:A01	ES 179	236965	1719.H12.gz43_219355
M00056044C:C01	ES 179	649900	1719.H15.gz43_219403
M00056044D:B04	ES 179	646579	1719.H19.gz43_219467
M00056044D:F01	ES 179	588502	1719.H22.gz43_219515
M00056044D:G07	ES 179	458683	1719.H24.gz43_219547
M00056045A:D03	ES 179	557676	1719.I01.gz43_219180
M00056045C:C09	ES 179	449814	1719.I10.gz43_219324
M00056045D:C09	ES 180	402476	1719.I21.gz43_219500
M00056045D:E06	ES 180	462293	1719.I23.gz43_219532
M00056046A:B04	ES 180	644738	1719.J02.gz43_219197
M00056046A;B12	ES 180	461517	1719.J04.gz43_219229
M00056046A:D12	ES 180	559550	1719.J06.gz43_219261
M00056046A:G08	ES 180	452102	1719.J07.gz43_219277
M00056046D:C11	ES 180	645070	1719.K04.gz43_219230
M00056047A;E09	ES 180	650348	1719.K09.gz43_219310
M00056047B:D11	ES 180	639711	1719.K17.gz43_219438
M00056047C:E03	ES 180	639255	1719,K22.gz43_219518
M00056047C:E08	ES 180	454844	1719.K23.gz43_219534
M00056047D:G10	ES 180	557853	1719.L06.gz43_219263
M00056048A:D12	ES 180	644894	1719.L09.gz43_219311
M00056048B:E01	ES 180	451544	1719.L12.gz43_219359
M00056048D:B09	ES 180	456840	1719.L19.gz43_219471
M00056049A;C11	ES 180	612572	1719.L22.gz43_219519
M00056049B:E11	ES 180	645942	1719.M04.gz43_219232
M00056049C:A08	ES 180	642184	1719.M06.gz43_219264
M00056049C:H04	ES 180	643748	1719.M11.gz43_219344
M00056049C:H07	ES 180	554070	1719.M12.gz43_219360
M00056049D:D03	ES 180	645431	1719.M16.gz43_219424
M00056049D:E01	ES 180	641715	1719.M17.gz43_219440

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CloneID	ES No	ClusterID	SequenceName
M00056050A:B05	ES 180	644686	1719.M19.gz43_219472
M00056050A:D02	ES 180	558439	1719.M21.gz43_219504
M00056050C:A09	ES 180	644639	1719.N01.gz43_219185
M00056050D:F04	ES 180	406734	1719.N07.gz43_219281
M00056051A;C03	ES 180	649259	1719.N11.gz43_219345
M00056051A:H11	ES 180	528616	1719.N12.gz43_219361
M00056051B;B03	ES 180	645375	1719.N15.gz43_219409
M00056051C;C09	ES 180	595066	1719.N21.gz43_219505
M00056051C:H09	ES 180	587106	1719.N23.gz43_219537
M00056051D:A07	ES 180	640603	1719.N24.gz43_219553
M00056051D:H02	ES 180	645151	1719.O05.gz43_219250
M00056052A:A11	ES 180	649429	1719.O06.gz43_219266
M00056052A;C07	ES 180	644819	1719.007.gz43_219282
M00056052C:G07	ES 180	647038	1719.016.gz43_219426
M00056052D:G01	ES 180	641786	1719.O22.gz43_219522
M00056053A:F01	ES 180	506920	1719.P03.gz43_219219
M00056053C:B04	ES 180	645814	1719.P08.gz43 219299
M00056053C:E04	ES 180	232093	1719.P10.gz43 219331
M00056053D:D07	ES 180	497101	1719.P15.gz43 219411
M00056054A:B06	ES 180	644692	1719.P17.gz43 219443
M00056054A:D09	ES 180	465589	1719.P18.gz43 219459
M00056054A:E03	ES 180	467255	1719.P19.gz43 219475
M00056054A:G07	ES 180	647306	1719.P22.gz43_219523
M00056054B:G05	ES 180	645049	1720.A02.gz43_219572
M00056054B:H11	ES 180	648013	1720.A03.gz43_219588
M00056054C:C09	ES 180	558494	1720.A04.gz43_219604
M00056054C:E12	ES 180	454906	1720,A06.gz43_219636
M00056055A:E04	ES 180	447676	1720.A14.gz43_219764
M00056055B;B06	ES 180	640818	1720,A18,gz43 219828
M00056057B:D01	ES 180	466795	1720.C02.gz43 219574
M00056057B:E12	ES 180	644927	1720,C04.gz43_219606
M00056057C:B02	ES 180	648688	1720.C05.gz43 219622
M00056057C:D06	ES 180	456059	1720.C08.gz43 219670
M00056057C:E01	ES 180	447635	1720,C09,gz43 219686
M00056057C:E12	ES 180	644928	1720,C12,gz43 219734
M00056057D:E11	ES 180	641469	1720.C15.gz43 219782
M00056058A:B07	ES 180	558154	1720,C16.gz43 219798
M00056058A;F08	ES 180	467780	1720.C19.gz43 219846
M00056058B:H09	ES 180	452630	1720,C22.gz43 219894
M00056059A:A02	ES 180	644597	1720,D06,gz43 219639
M00056059A:B01	ES 180	649054	1720.D08.gz43 219671
M00056059B:D10	ES 180	473007	1720,D11.gz43 219719
M00056060C:C06	ES 180	448627	1720,E04.gz43 219608
M00056060D:H11	ES 180	642166	1720,E10.gz43 219704

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00056061A:E05	ES 180	644916	1720.E12.gz43_219736
M00056061B:A03	ES 180	641801	1720.E13.gz43_219752
M00056062D:D08	ES 180	446415	1720.F20.gz43_219865
M00056062D:F12	ES 180	643510	1720.G01.gz43_219562
M00056063C:F10	ES 180	375577	1720.G15.gz43_219786
M00056063C:G11	ES 180	650549	1720.G16.gz43_219802
M00056065B:A02	ES 180	641253	1720.H06.gz43_219643
M00056065D:C11	ES 180	643381	1720.H16.gz43_219803
M00056066A;D03	ES 180	553923	1720.H20.gz43_219867
M00056066B:H04	ES 180	14157	1720,I02.gz43_219580
M00056066D:G04	ES 180	461351	1720.I16.gz43_219804
M00056067A:A12	ES 180	455716	1720.I18.gz43_219836
M00056067B:D08	ES 180	645900	1720.I21.gz43_219884
M00056068D:A06	ES 180	644822	1720.J18.gz43_219837
M00056068D:E06	ES 180	641253	1720.J22.gz43_219901
M00056069A;F01	ES 180	460047	1720.K06.gz43_219646
M00056069A:F06	ES 180	644225	1720.K08.gz43_219678
M00056069B:E04	ES 180	645076	1720.K14.gz43_219774
M00056070C:B09	ES 180	644701	1720.L12.gz43_219743
M00056072B:C06	ES 180	551181	1720.M11.gz43_219728
M00056072D:E08	ES 180	649364	1720.M21.gz43_219888
M00056073B:H11	ES 180	636654	1720.N08.gz43_219681
M00056074B:D12	ES 180	643348	1720.N23.gz43_219921
M00056074B:E11	ES 180	641135	1720.O02.gz43_219586
M00056074C:H08	ES 180	487567	1720.O12.gz43_219746
M00056074C:H09	ES 180	651108	1720.O13.gz43_219762
M00056077B:E03	ES 180	448485	1729.A06.gz43_217713
M00056077C:H02	ES 180	646696	1729.A12.gz43_217809
M00056078A:D07	ES 180	479868	1729.A20.gz43_217937
M00056078B:G02	ES 180	650900	1729.B04.gz43_217682
M00056079A:B01	ES 180	613722	1729.B18.gz43_217906
M00056080C:D04	ES 180	464778	1729,C22.gz43_217971
M00056081A:C05	ES 180	649149	1729.D04.gz43_217684
M00056081B:F09	ES 180	640891	1729.D12.gz43_217812
M00056082A:E12	ES 180	560054	1729.E11.gz43_217797
M00056082B:G07	ES 180	570939	1729.E15.gz43_217861
M00056083D:D11	ES 180	647437	1729.F16.gz43_217878
M00056084C:D11	ES 180	649202	1729.G03.gz43_217671
M00056087B:G12	ES 180	644715	1729.H12.gz43_217816
M00056087D:E04	ES 180	446243	1729.H22.gz43_217976
M00056087D:G07	ES 180	649520	1729.H24.gz43_218008
M00056089D:A09	ES 180	454176	1729.J05.gz43_217706
M00056089D:E07	ES 180	649356	1729.J08.gz43_217754
M00056091A:E04	ES 180	644054	1729.K04.gz43_217691
M00056089D:A09 M00056089D:E07	ES 180 ES 180	454176 649356	1729.H24.gz43_21800 1729.J05.gz43_21770 1729.J08.gz43_21775

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00056091C;C06	ES 180	644712	1729.K11.gz43_217803
M00056091C:D09	ES 180	645050	1729.K13.gz43_217835
M00056091C:H03	ES 180	464267	1729.K15.gz43_217867
M00056091D:C12	ES 180	645409	1729.K17.gz43_217899
M00056091D:E10	ES 180	649328	1729.K20.gz43_217947
M00056092B:D10	ES 180	535436	1729.L02.gz43_217660
M00056092C:E12	ES 180	710155	1729.L08.gz43_217756
M00056093A;B12	ES 180	650547	1729.L16.gz43_217884
M00056093A.F08	ES 180	418763	1729.L18.gz43_217916
M00056093B:D03	ES 180	645131	1729.L20.gz43_217948
M00056095A:C02	ES 180	644543	1729.M22.gz43_217981
M00056095B:A07	ES 180	643440	1729.N06.gz43_217726
M00056095C:E02	ES 180	644965	1729.N12.gz43_217822
M00056095C:G07	ES 180	649514	1729.N14.gz43_217854
M00056097B:C04	ES 180	649082	1729.P04.gz43_217696
M00056097B:G01	ES 180	649570	1729.P06.gz43_217728
M00056098B:C04	ES 180	649054	1729.P16.gz43_217888
M00056098D:A08	ES 180	597647	1729.P22.gz43_217984
M00056120C:H04	ES 180	648654	1731,A01.gz43_219940
M00056120D:F01	ES 180	644037	1731.A02.gz43_219956
M00056121A;E05	ES 180	464990	1731.A09.gz43_220068
M00056121D:A12	ES 180	642204	1731.A19.gz43_220228
M00056121D:C11	ES 180	648013	1731.A20.gz43_220244
M00056122A:A05	ES 180	503122	1731.A22.gz43_220276
M00056122A:B05	ES 180	589098	1731.A23.gz43_220292
M00056122A:D02	ES 180	648076	1731.A24.gz43_220308
M00056122B:A07	ES 180	639555	1731.B04.gz43_219989
M00056122B:G01	ES 180	648109	1731.B07.gz43_220037
M00056122B:G09	ES 180	627515	1731.B08.gz43_220053
M00056122C:H12	ES 180	460190	1731.B12.gz43_220117
M00056122D:B07	ES 180	557419	1731.B13.gz43_220133
M00056122D:F10	ES 180	646917	1731.B17.gz43_220197
M00056123B:F02	ES 180	531461	1731.B23.gz43_220293
M00056123B:G05	ES 180	648547	1731.C01.gz43_219942
M00056123D:A07	ES 180	627341	1731.C05.gz43_220006
M00056124B:C12	ES 180	477555	1731.C14.gz43_220150
M00056124D:F06	ES 180	515350	1731.C22.gz43_220278
M00056124D:G03	ES 180	455808	1731.C23.gz43_220294
M00056125B:F01	ES 180	465127	1731.D04.gz43_219991
M00056127A:A10	ES 180	422242	1731.D07.gz43_220039
M00056127A:H03	ES 180	644928	1731.D11.gz43_220103
M00056127B:E11	ES 180	638934	1731.D13.gz43_220135
M00056127C:C02	ES 180	639774	1731.D15.gz43_220167
M00056127C:C06	ES 180	649095	1731.D16.gz43_220183

Table 13

CloneID	ES No	ClusterID	SequenceName
M00056127D:G10	ES 180	643510	1731.D23.gz43_220295
M00056128B:A07	ES 180	456985	1731,E07.gz43_220040
M00056128B:D03	ES 180	452941	1731.E08.gz43_220056
M00056128C:B11	ES 180	646584	1731.E11.gz43_220104
M00056128C:F02	ES 180	647248	1731.E14.gz43_220152
M00056129A:G01	ES 180	647431	1731.E21.gz43_220264
M00056129C:F09	ES 180	476380	1731.F03.gz43_219977
M00056130B:E11	ES 180	646785	1731.F13.gz43_220137
M00056130C:C12	ES 180	649299	1731.F15.gz43_220169
M00056130C:F08	ES 180	647290	1731.F16.gz43_220185
M00056130D:E06	ES 180	556336	1731.F22.gz43_220281
M00056131A:E09	ES 180	453726	1731.G05.gz43_220010
M00056131B:E01	ES 180	562378	1731.G12.gz43_220122
M00056131C:H03	ES 180	649735	1731.G16.gz43_220186
M00056131D:B02	ES 180	503628	1731.G19.gz43_220234
M00056132B:C12	ES 180	635354	1731.H02.gz43_219963
M00056132C:F04	ES 180	557026	1731.H06.gz43_220027
M00056133B:F01	ES 180	642936	1731.H18.gz43_220219
M00056133D:D09	ES 180	463824	1731.I01.gz43_219948
M00056134A:D04	ES 180	640695	1731.I07.gz43_220044
M00056134B:B10	ES 180	640737	1731.I10.gz43_220092
M00056134C:C06	ES 180	470006	1731 I14.gz43_220156
M00056134D:B07	ES 180	708175	1731.I21.gz43_220268
M00056134D:F02	ES 180	649482	1731.I24.gz43_220316
M00056134D:G07	ES 180	711493	1731.J02.gz43_219965
M00056134D:G11	ES 180	649617	1731.J03.gz43_219981
M00056135B:E03	ES 180	459521	1731.J10.gz43_220093
M00056135C:C05	ES 180	646039	1731.J16.gz43_220189
M00056135C:C08	ES 180	642650	1731.J17.gz43_220205
M00056135C:H06	ES 180	557867	1731.J19.gz43_220237
M00056136A;B11	ES 180	648996	1731.J23.gz43_220301
M00056136D:B08	ES 180	641957	1731.K06.gz43_220030
M00056137B:F07	ES 180	648311	1731.K14.gz43_220158
M00056137B:F11	ES 180	478393	1731.K15.gz43_220174
M00056137C:A01	ES 180	642275	1731.K17.gz43_220206
M00056137C:E01	ES 180	639395	1731.K19.gz43_220238
M00056137C:G02	ES 180	645367	1731.K20.gz43_220254
M00056137D:H05	ES 181	648742	1731.L02.gz43_219967
M00056138A:B11	ES 181	458938	1731.L04.gz43_219999
M00056138B:F05	ES 181	549691	1731.L10.gz43_220095
M00056138D:G08	ES 181	651015	1731.L17.gz43_220207
M00056139A:D12	ES 181	647232	1731.L20.gz43_220255
M00056139D:E05	ES 181	648341	1731.M08.gz43_220064

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00056139D:H04	ES 181	514838	1731.M10.gz43_220096
M00056140A:A07	ES 181	450049	1731.M11.gz43_220112
M00056140A:F12	ES 181	646581	1731.M16.gz43_220192
M00056140B:G06	ES 181	649557	1731.M22.gz43_220288
M00056140C:E04	ES 181	648721	1731,N01.gz43_219953
M00056140D:G07	ES 181	649278	1731.N07.gz43_220049
M00056141C:C10	ES 181	650745	1731.N13.gz43_220145
M00056141C:H01	ES 181	470667	1731.N16.gz43_220193
M00056141C:H07	ES 181	644686	1731.N17.gz43_220209
M00056141D:D09	ES 181	562755	1731.N22.gz43_220289
M00056141D:E08	ES 181	446164	1731.N23.gz43_220305
M00056142A:F03	ES 181	648481	1731.O01.gz43_219954
M00056142B:D02	ES 181	650076	1731.O06.gz43_220034
M00056142C:A09	ES 181	452738	1731.O10.gz43_220098
M00056143A:H08	ES 181	419114	1731.O23.gz43_220306
M00056144B:C01	ES 181	558377	1731.P07.gz43_220051
M00056144D:C05	ES 181	647949	1731.P12.gz43_220131
M00056146A:A09	ES 181	550714	1732.A05.gz43_220388
M00056146A:H09	ES 181	648667	1732.A12.gz43_220500
M00056146B:E05	ES 181	640634	1732.A14.gz43_220532
M00056146D:D04	ES 181	488592	1732.A20.gz43_220628
M00056149C:A02	ES 181	465734	1732.C11.gz43_220486
M00056149C:B01	ES 181	459012	1732.C12.gz43_220502
M00056149C:E10	ES 181	650877	1732.C16.gz43_220566
M00056149D:F06	ES 181	639485	1732.C22.gz43_220662
M00056150A:E04	ES 181	650820	1732.C24.gz43_220694
M00056151A:E07	ES 181	463513	1732.D16.gz43_220567
M00056152A:B11	ES 181	605761	1732.E11.gz43_220488
M00056152A:D07	ES 181	461135	1732.E13.gz43_220520
M00056152C:G08	ES 181	651075	1732.E20.gz43_220632
M00056153A:G04	ES 181	644221	1732.F07.gz43_220425
M00056153A:H07	ES 181	644801	1732.F09.gz43_220457
M00056153B:F11	ES 181	449882	1732.F16.gz43_220569
M00056154B:D02	ES 181	649443	1732.G08.gz43_220442
M00056154B:F11	ES 181	650249	1732.G10.gz43_220474
M00056155A:G09	ES 181	640162	1732.G24.gz43_220698
M00056155B:A02	ES 181	646323	1732.H01.gz43_220331
M00056157A:B11	ES 181	416624	1732.I08.gz43_220444
M00056157B:A08	ES 181	642244	1732.I12.gz43_220508
M00056157B:H06	ES 181	446184	1732.I14.gz43_220540
M00056157C:E10	ES 181	554581	1732.I19.gz43_220620
M00056157D:C02	ES 181	648930	1732.I21.gz43_220652
M00056157D:H08	ES 181	646596	1732.I23.gz43_220684
M00056158C:D11	ES 181	649482	1732.J05.gz43_220397

Table 13

Table 13	1		
CloneID	ES No	ClusterID	SequenceName
M00056158C:F09	ES 181	650987	1732.J06.gz43_220413
M00056159A:A08	ES 181	644856	1732.J14.gz43_220541
M00056159A:E03	ES 181	650909	1732.J16.gz43_220573
M00056159B;E06	ES 181	648774	1732.J21.gz43_220653
M00056159C:C11	ES 181	459260	1732.J23.gz43_220685
M00056160B:D06	ES 181	414821	1732.K09.gz43_220462
M00056160C:F12	ES 181	463966	1732.K15.gz43_220558
M00056160C:G01	ES 181	645100	1732.K16.gz43_220574
M00056160D:C06	ES 181	649991	1732.K20.gz43_220638
M00056161C:G06	ES 181	650348	1732.L12.gz43_220511
M00056161C:H10	ES 181	468262	1732.L13.gz43_220527
M00056162A:B06	ES 181	467972	1732.L19.gz43_220623
M00056162A:C09	ES 181	649947	1732.L20.gz43_220639
M00056162A:E09	ES 181	555634	1732.L21.gz43_220655
M00056162A:F01	ES 181	634409	1732.L23.gz43_220687
M00056162A:G09	ES 181	584499	1732.L24.gz43_220703
M00056162C:F02	ES 181	647412	1732.M07.gz43_220432
M00056162D:A01	ES 181	649782	1732.M11.gz43_220496
M00056162D:D03	ES 181	650116	1732.M13.gz43_220528
M00056162D:D06	ES 181	380477	1732.M14.gz43_220544
M00056163A:B10	ES 181	649883	1732.M17.gz43_220592
M00056164A:H03	ES 181	224092	1732.N20.gz43_220641
M00056165C:B11	ES 181	650534	1732.O13.gz43_220530
M00056165D:D09	ES 181	649578	1732.O15.gz43_220562
M00056167C:H06	ES 181	414006	1732.P18.gz43_220611
M00056168A:C06	ES 181	558915	1741.A01.gz43_220708
M00056168C:A05	ES 181	649795	1741.A09.gz43_220836
M00056169B:C02	ES 181	458683	1741.A21.gz43_221028
M00056169C:F07	ES 181	458085	1741.B05.gz43_220773
M00056169D:A03	ES 181	555771	1741,B08,gz43 220821
M00056169D:D05	ES 181	648777	1741.B12.gz43 220885
M00056169D:H05	ES 181	650800	1741.B16.gz43_220949
M00056170B:B09	ES 181	642256	1741.B24.gz43_221077
M00056170B:C04	ES 181	602324	1741.C01.gz43 220710
M00056170C:C09	ES 181	394772	1741.C06.gz43 220790
M00056171A:H01	ES 181	646173	1741.C20.gz43 221014
M00056171C:A08	ES 181	650063	1741.D02.gz43 220727
M00056172A:H06	ES 181	561747	1741.D14.gz43 220919
M00056172B:A12	ES 181	648855	1741.D15.gz43 220935
M00056173D:C05	ES 181	646699	1741.E07.gz43_220808
M00056174B:D02	ES 181	646915	1741.E14.gz43_220920
M00056174B:E02	ES 181	647086	1741.E16.gz43_220952
M00056174B:E04	ES 181	640285	1741.E17.gz43_220968
M00056174B:H11	ES 181	557010	1741.E20.gz43_221016

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CloneID	ES No	ClusterID	SequenceName
M00056174C:F06	ES 181	479997	1741.E24.gz43_221080
M00056174C:H09	ES 181	648063	1741.F04.gz43_220761
M00056175A:A06	ES 181	453768	1741.F10.gz43_220857
M00056175A:C04	ES 181	649136	1741.F11.gz43 220873
M00056175C:A10	ES 181	645288	1741.F19.gz43 221001
M00056175C:B11	ES 181	650146	1741.F20.gz43 221017
M00056175D:A06	ES 181	648820	1741.F24.gz43 221081
M00056175D:E05	ES 181	649299	1741.G03.gz43 220746
M00056176A:D06	ES 181	593173	1741.G07.gz43_220810
M00056176B:B05	ES 181	457272	1741.G10.gz43_220858
M00056176B:E10	ES 181	446289	1741.G12.gz43_220890
M00056176C:B11	ES 181	648865	1741.G14.gz43_220922
M00056177A:C10	ES 181	140459	1741.G19.gz43_221002
M00056177A:D12	ES 181	499690	1741.G20.gz43_221018
M00056177A:H01	ES 181	610986	1741.G23.gz43_221066
M00056177C:E08	ES 181	639996	1741.H06.gz43_220795
M00056178B:A07	ES 181	642557	1741.H14.gz43 220923
M00056178B:A11	ES 181	638730	1741.H15.gz43 220939
M00056178B:H08	ES 181	459923	1741.H17.gz43 220971
M00056178C:C02	ES 181	464778	1741.H18.gz43 220987
M00056178C:E12	ES 181	584745	1741.H20.gz43_221019
M00056178C:F02	ES 181	640922	1741.H21.gz43_221035
M00056179A:D12	ES 181	379805	1741,I08.gz43_220828
M00056179B:H12	ES 181	639886	1741,I14.gz43_220924
M00056180B:A06	ES 181	644053	1741,I23.gz43_221068
M00056180B:D05	ES 181	492982	1741,J02.gz43_220733
M00056180B:F11	ES 181	641193	1741.J04.gz43_220765
M00056180D:E06	ES 181	378694	1741.J10.gz43_220861
M00056180D:F05	ES 181	449452	1741.J12.gz43_220893
M00056180D:G07	ES 181	183842	1741.J14.gz43_220925
M00056181A:B05	ES 181	471836	1741.J16.gz43_220957
M00056181A:E07	ES 181	452729	1741.J17.gz43_220973
M00056181B:B02	ES 181	640514	1741.J19.gz43_221005
M00056181B:D06	ES 181	645521	1741.J20.gz43_221021
M00056181B:E11	ES 181	561793	1741.J21.gz43_221037
M00056181C:D02	ES 181	645578	1741.J22.gz43_221053
M00056182A:B04	ES 181	557353	1741.K02.gz43_220734
M00056182A:G11	ES 181	643206	1741.K07.gz43_220814
M00056182B:G12	ES 181	460506	1741.K09.gz43_220846
M00056182C:F08	ES 181	469409	1741.K14.gz43_220926
M00056182D:A07	ES 181	641525	1741.K15.gz43_220942
M00056183A:B09	ES 181	459974	1741.K19.gz43_221006
M00056183B:G11	ES 181	561856	1741.L04.gz43_220767
M00056183C:F01	ES 181	710362	1741.L08.gz43_220831

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M00056183C:F11	Table 13			
M00056184A:B11   ES 181   649702   1741.L14.gzt3 220927   M00056184B:D08   ES 181   463487   1741.L18.gzt3 220927   M00056184B:H08   ES 181   4646987   1741.L21.gzt3 220927   M00056184C:A09   ES 181   647688   1741.L22.gzt3 221055   M00056184C:A09   ES 181   667688   1741.L22.gzt3 221055   M00056184C:C03   ES 181   667688   1741.L22.gzt3 221055   M00056184C:C03   ES 181   662631   1741.M04.gzt3 220768   M00056184C:E08   ES 181   642631   1741.M04.gzt3 220768   M00056184C:E08   ES 181   642631   1741.M04.gzt3 220768   M00056184C:E08   ES 181   642631   1741.M07.gzt3 220816   M00056184D:D06   ES 181   655555   1741.M12.gzt3 220849   M00056185D:D06   ES 181   639132   1741.M12.gzt3 220849   M00056185D:C01   ES 181   653023   1741.M07.gzt3 220818   M00056185D:C06   ES 181   649958   1741.M09.gzt3 220849   M00056185D:D06   ES 181   649958   1741.M09.gzt3 220849   M00056185D:D06   ES 181   649957   1741.M12.gzt3 220849   M00056185D:D06   ES 181   649957   1741.M12.gzt3 220849   M00056185D:D06   ES 181   649958   1741.M09.gzt3 220849   M00056185D:D06   ES 181   644801   1741.M2.gzt3 220849   M00056185D:D06   ES 181   644801   1741.M2.gzt3 220849   M00056186D:D09   ES 181   644801   1741.M2.gzt3 220849   M00056186D:D09   ES 181   644801   1741.M2.gzt3 220849   M00056186D:D09   ES 181   644801   1741.M2.gzt3 220849   M00056186D:G05   ES 181   644801   1741.M2.gzt3 220849   M00056186D:G05   ES 181   644801   1741.D012.gzt3 220849   M00056186D:G05   ES 181   644801   1741.D012.gzt3 220849   M00056186D:G05   ES 181   644801   1741.D012.gzt3 220849   M00056186D:G05   ES 181   644801   1741.D012.gzt3 220849   M00056186D:G05   ES 181   644801   1741.D012.gzt3 220849   M00056186D:G05   ES 181   644801   1741.D012.gzt3 220849   M00056186D:G05   ES 181   644801   1741.D012.gzt3 220849   M00056186D:G05   ES 181   644801   1741.D012.gzt3 220849   M00056186D:G05   ES 181   644801   1741.D012.gzt3 220849   M00056186D:G05   ES 181   644801   1741.D012.gzt3 220849   M00056186D:G05   ES 181   644801   1741.D012.gzt3 220849   M0005618	CloneID	ES No	ClusterID	SequenceName
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M00056184B:H08				
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M00056184C:C07				
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M00056188E:D04	M00056187B:H02	ES 181	644819	1741.O12.gz43_220898
M00056188B:E07   ES 181   648344   1741.P04.gz43 220771	M00056188A:E05	ES 181	641945	1741.O21.gz43_221042
M00056188B:E12   ES 181   639520   1741.P05_gz43_220787   M00056188C:D02   ES 181   650188   1741.P08_gz43_220832   M00056188C:G06   ES 181   640443   1741.P11_gz43_220883   M00056188C:H04   ES 181   648665   1741.P12_gz43_220883   M00056189C:H04   ES 181   562152   1741.P18_gz43_220893   M00056189B:A01   ES 181   562152   1741.P18_gz43_220983   M00056189B:A01   ES 181   465594   1741.P19_gz43_221011   M00056191A:H04   ES 181   465594   1742.P08_gz43_221141   M00056191A:H04   ES 181   646499   1742.P08_gz43_221141   M00056192A:D01   ES 181   645560   1742.B08_gz43_221238   M00056195C:F12   ES 181   644916   1742.E08_gz43_221273   M00056195C:D07   ES 181   649402   1742.E08_gz43_221273   M00056196C:B03   ES 181   397363   1742.F08_gz43_22125   M00056197C:H01   ES 181   640101   1742.F08_gz43_22125   M00056197C:H01   ES 181   640101   1742.F08_gz43_22125   M00056197C:H01   ES 181   640101   1742.F08_gz43_221240   M00056197C:H01   ES 181   640913   1742.F08_gz43_221240   M00056197C:H01   ES 181   640913   1742.F08_gz43_221240   M00056203D:F05   ES 181   640913   1742.I08_gz43_221240   M00056203D:F05   ES 181   64368   1742.J08_gz43_22123   M00056203D:F05   ES 181   570939   1742.I08_gz43_221240   M00056203D:H03   ES 181   570939   1742.I08_gz43_221149	M00056188B:D04	ES 181	642164	1741.P01.gz43_220723
M00056188C:D02	M00056188B:E07	ES 181	648344	1741.P04.gz43 220771
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M00056188C:H04	M00056188C:D02	ES 181	650188	1741.P08.gz43_220835
M00056189A:H09	M00056188C:G06	ES 181	640443	1741.P11.gz43_220883
M00056189E:A01   ES 181   465594   1741.P19.gz43_221011     M00056191A:H04   ES 181   646499   1742.B08.gz43_221014     M00056191B:E08   ES 181   645762   1742.B08.gz43_221145     M00056192A:D01   ES 181   645560   1742.B19.gz43_22125     M00056195C:F12   ES 181   644916   1742.E08.gz43_22137     M00056195D:D07   ES 181   644916   1742.E08.gz43_22137     M00056196C:B03   ES 181   397363   1742.P03.gz43_22137     M00056196C:G12   ES 181   648590   1742.F03.gz43_22122     M00056196C:F101   ES 181   648590   1742.P03.gz43_221240     M00056196A:B05   ES 181   649011   1742.P20.gz43_22149     M00056196A:B05   ES 181   649011   1742.P20.gz43_221240     M0005620D:D05   ES 181   649013   1742.P03.gz43_221410     M0005620D:F05   ES 181   649013   1742.103.gz43_22124     M0005620D:F05   ES 181   649013   1742.103.gz43_22123     M0005620D:F05   ES 181   649013   1742.103.gz43_22124     M0005620D:F05   ES 181   649013   1742.103.gz43_22123     M0005620D:F05   ES 181   649013   1742.103.gz43_22123     M0005620D:F05   ES 181   649013   1742.103.gz43_22123     M0005620D:F05   ES 181   649013   1742.103.gz43_22123     M0005620D:F05   ES 181   649013   1742.103.gz43_22123     M0005620D:F05   ES 181   649013   1742.103.gz43_22123     M0005620D:F05   ES 181   649013   1742.203.gz43_22123     M0005620D:F05   ES 181   649013   1742.203.gz43_22123     M0005620D:F05   ES 181   649013   1742.203.gz43_22134     M0005620D:F05   ES 181   649013   1742.203.gz43_22134     M0005620D:F05   ES 181   649013   1742.203.gz43_22134     M0005620D:F05   ES 181   649013   1742.203.gz43_22134     M0005620D:F05   ES 181   649013   1742.203.gz43_22134     M0005620D:F05   ES 181   649013   1742.203.gz43_22134     M0005620D:F05   ES 181   649013   1742.203.gz43_22134     M0005620D:F05   ES 181   649013   1742.203.gz43_22134     M0005620D:F05   ES 181   649013   1742.203.gz43_22134     M0005620D:F05   ES 181   649013   1742.203.gz43_22134     M0005620D:F05   ES 181   649013   1742.203.gz43_22134     M0005620D:F05   ES 181   649013   1742.203.gz43_22134	M00056188C:H04	ES 181	648665	1741.P12.gz43 220899
M00056191A:H04	M00056189A:H09	ES 181	562152	1741.P18.gz43_220995
M00056191B:E08   ES 181   453762   1742.B08_gz43_221205   M00056192A:D01   ES 181   645560   1742.B19_gz43_221381   M00056192A:D02   ES 181   644916   1742.B19_gz43_221376   M00056195D:D07   ES 181   644916   1742.E12_gz43_22172   M00056196C:B03   ES 181   397363   1742.F03_gz43_2211276   M00056196C:B03   ES 181   397363   1742.F03_gz43_221129   M00056196C:G12   ES 181   648590   1742.F03_gz43_221222   M00056197C:H01   ES 181   640101   1742_F02_gz43_221240   M00056198A:B05   ES 181   647387   1742_G01_gz43_221249   M00056200D:D05   ES 181   452066   1742_H21_gz43_221419   M00056200D:D05   ES 181   640913   1742_103_gz43_22123   M00056203D:F05   ES 181   643168   1742_J03_gz43_22134   M00056203D:F05   ES 181   463168   1742_J03_gz43_221138   M00056203D:H03   ES 181   570939   1742_J04_gz43_22149   M00056203D:H03   ES 181   570939   1742_J04_gz43_221149   M00056203D:H03   ES 181   4566200   1742_J03_gz43_221149   M00056203D:H03   ES 181   570939   1742_J04_gz43_221449   M00056203D:H03   ES 181   4566200   1742_J03_gz43_221149   M00056204A:B08   ES 181   4566200   1742_J04_gz43_221449   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221149   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221135   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221149   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221149   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221149   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221149   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221149   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221149   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221149   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221149   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221149   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221149   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221169   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221169   M00056204A:B08   ES 181   4566200   1742_J03_gz43_21169   M00056204A:B08   ES 181   4566200   1742_J03_gz43_221169	M00056189B:A01	ES 181	465594	1741.P19.gz43_221011
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M00056195C:F12	M00056191B:E08	ES 181	453762	1742.B08.gz43_221205
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	M00056204A;C04	ES 181	647991	1742.J07.gz43 221197

Table 13

Table 13			
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M00056205D:G04	ES 181	650756	1742.K15.gz43_221326
M00056206B:A10	ES 181	645177	1742.K19.gz43 221390
M00056206D:E06	ES 181	639158	1742,L03.gz43 221135
M00056206D:F07	ES 181	463545	1742.L04.gz43 221151
M00056207C:B04	ES 181	446621	1742.L17.gz43 221359
M00056208B:C10	ES 181	172843	1742.M03.gz43 221136
M00056208C:A10	ES 181	559955	1742.M06.gz43_221184
M00056208D:A12	ES 181	645223	1742.M11.gz43_221264
M00042346A:B04	ES 182	448412	1766.A01.gz43_224099
M00042346B:D12	ES 182	448491	1766.A05.gz43_224163
M00042346C:G12	ES 182	448892	1766.A09.gz43_224227
M00042432D:G02	ES 182	485237	1766.E01.gz43_224103
M00042433C:H05	ES 182	477797	1766,E13,gz43_224295
M00042434B:F04	ES 182	450334	1766.E20.gz43_224407
M00042437B:A11	ES 182	479131	1766.G01.gz43_224105
M00042438C:H11	ES 182	451183	1766.G20.gz43_224409
M00042439A:D09	ES 182	481457	1766.H03.gz43_224138
M00042461C:E09	ES 182	483219	1766.I13.gz43_224299
M00042462A:F12	ES 182	450212	1766.I19.gz43_224395
M00042464A:F06	ES 182	451793	1766.J21.gz43_224428
M00042464C:B06	ES 182	448098	1766.K04.gz43_224157
M00042470B:H11	ES 182	447377	1766.M22.gz43_224447
M00042515B:A10	ES 182	449104	1767.A01.gz43_224483
M00042515D:B11	ES 182	449215	1767.A04.gz43_224531
M00042515D:G01	ES 182	451982	1767.A05.gz43_224547
M00056209B:D02	ES 182	507050	1742.M19.gz43_221392
M00056210B:C11	ES 182	470801	1742.N09.gz43_221233
M00056210B:E03	ES 182	649354	1742.N12.gz43_221281
M00056210D:H05	ES 182	457539	1742.N15.gz43_221329
M00056211A:C08	ES 182	648036	1742.N18.gz43_221377
M00056212C:C09	ES 182	642118	1742.P01.gz43_221107
M00056213C:C01	ES 182	646275	1742.P19.gz43_221395
M00056213C:G11	ES 182	466418	1742.P24.gz43_221475
M00056214B:H03	ES 182	638807	1743.A08.gz43_221588
M00056214C:A04	ES 182	499517	1743.A10.gz43_221620
M00056214D:F05	ES 182	650975	1743.A16.gz43_221716
M00056216B:H09	ES 182	648953	1743.B17.gz43_221733
M00056217A:H11	ES 182	648155	1743.C04.gz43_221526
M00056217B:A06	ES 182	650487	1743.C05.gz43_221542
M00056217B:G03	ES 182	562359	1743.C09.gz43_221606
M00056217C:B12	ES 182	602324	1743.C10.gz43_221622

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M00056218C:C02	ES 182	648351	1743.D05.gz43 221543
M00056218D:H06	ES 182	411603	1743.D14.gz43 221687
M00056220A:H04	ES 182	644916	1743.E16.gz43_221720
M00056220B:F11	ES 182	722253	1743.F01.gz43 221481
M00056222A:E05	ES 182	570052	1743.G11.gz43 221642
M00056222D:C06	ES 182	574166	1743.G18.gz43_221754
M00056223A;B11	ES 182	648989	1743.G23.gz43_221834
M00056223A;C03	ES 182	442347	1743.G24.gz43_221850
M00056223A:D10	ES 182	574166	1743.H01.gz43_221483
M00056223A:H07	ES 182	649709	1743.H03.gz43_221515
M00056223C:C07	ES 182	408428	1743.H11.gz43_221643
M00056224B:F01	ES 182	645549	1743.H17.gz43_221739
M00056224D:E08	ES 182	453470	1743.H23.gz43_221835
M00056225A:D08	ES 182	556635	1743.I01.gz43_221484
M00056225B:A11	ES 182	618670	1743.I06.gz43_221564
M00056227C:A01	ES 182	648777	1743.J20.gz43_221789
M00056227D;A02	ES 182	578718	1743.K04.gz43_221534
M00056228A:A02	ES 182	447892	1743.K09.gz43_221614
M00056228C:B04	ES 182	465536	1743.K24.gz43_221854
M00056229C:F05	ES 182	33738	1743.L16.gz43_221727
M00056231B:E01	ES 182	649351	1743.M11.gz43_221648
M00056232B;A11	ES 182	451487	1743.M21.gz43_221808
M00056232B:C04	ES 182	649030	1743.M23.gz43_221840
M00056232C:E06	ES 182	649272	1743.N04.gz43_221537
M00056232D:C08	ES 182	502413	1743.N09.gz43_221617
M00056235A:C12	ES 182	648816	1743.O22.gz43_221826
M00056236A:G12	ES 182	706245	1743.P07.gz43_221587
M00056236B:B03	ES 182	636876	1743.P09.gz43_221619
M00056237B:D04	ES 182	703978	1744,A06.gz43_221940
M00056237C:C05	ES 182	470292	1744.A10.gz43_222004
M00056238A;A03	ES 182	707483	1744.A19.gz43_222148
M00056238C:C11	ES 182	649042	1744.B08.gz43_221973
M00056239A:D06	ES 182	648159	1744.B14.gz43_222069
M00056240B:F08	ES 182	649396	1744.C10.gz43_222006
M00056240D:H07	ES 182	447272	1744.C15.gz43_222086
M00056242D:A02	ES 182	648989	1744.E04.gz43_221912
M00056243C:G10	ES 182	651073	1744.E22.gz43_222200
M00056244A;A01	ES 182	650463	1744.F03.gz43_221897
M00056244A;B01	ES 182	650549	1744.F04.gz43_221913
M00056245A:D11	ES 182	650463	1744.F24.gz43_222233
M00056246A:E01	ES 182	637387	1744.G17.gz43_222122
M00056246C:G07	ES 182	419255	1744.H08.gz43_221979

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T HISTO TO			
CloneID	ES No	ClusterID	SequenceName
M00056246D:A07	ES 182	549178	1744.H09.gz43_221995
M00056247A:D02	ES 182	162981	1744.H17.gz43_222123
M00056247A:F07	ES 182	457221	1744.H18.gz43_222139
M00056247B:C11	ES 182	650018	1744.H21.gz43_222187
M00056249A:F11	ES 182	650919	1744.I24.gz43_222236
M00056249A:H07	ES 182	645872	1744.J02.gz43_221885
M00056249B:C02	ES 182	506719	1744,J03.gz43_221901
M00056249C:E06	ES 182	650860	1744.J10.gz43_222013
M00056250C:A08	ES 182	408961	1744.K04.gz43_221918
M00056250C:D05	ES 182	650661	1744.K07.gz43_221966
M00056251B;A12	ES 182	643327	1744.K14.gz43_222078
M00056251B:B07	ES 182	452646	1744.K15.gz43_222094
M00056251C:H04	ES 182	586794	1744.K24.gz43_222238
M00056253A:F12	ES 182	648996	1744.M08.gz43_221984
M00056253B:H04	ES 182	639119	1744.M14.gz43_222080
M00056253C:C04	ES 182	703217	1744,M18.gz43_222144
M00056253D:G04	ES 182	641874	1744.M23.gz43_222224
M00056254A:F04	ES 182	651051	1744.N02.gz43_221889
M00056254D:C04	ES 182	451361	1744.N09.gz43_222001
M00056255B;F09	ES 182	559324	1744,N18.gz43_222145
M00056255C:C04	ES 182	645472	1744.N20.gz43_222177
M00056256A:C02	ES 182	645271	1744.O01.gz43_221874
M00056256C:D12	ES 182	643944	1744.O17.gz43_222130
M00056258A:A11	ES 182	464091	1744.P19.gz43_222163
M00056258A:E10	ES 182	610269	1744.P23.gz43_222227
M00056258C:D07	ES 182	633594	1753.A05.gz43_222308
M00056258D:H06	ES 182	648590	1753.A12.gz43_222420
M00056259A:E08	ES 182	555021	1753.A17.gz43_222500
M00056259C:E03	ES 182	459914	1753.A24.gz43_222612
M00056260C:E11	ES 182	460245	1753.B14.gz43_222453
M00056260C:F08	ES 182	552839	1753.B17.gz43_222501
M00056261A:F12	ES 182	645271	1753.C03.gz43_222278
M00056262A:B07	ES 182	647897	1753.C18.gz43_222518
M00056262B:B08	ES 182	403111	1753.C23.gz43_222598
M00056262D:B11	ES 182	446572	1753.D04.gz43_222295
M00056262D:G08	ES 182	648534	1753.D06.gz43_222327
M00056263A:D08	ES 182	649262	1753.D10.gz43_222391
M00056263B:E10	ES 182	486452	1753.D17.gz43_222503
M00056264A:G05	ES 182	650679	1753.E03.gz43_222280
M00056264B:C03	ES 182	648063	1753,E06.gz43_222328
M00056265A;E01	ES 182	646609	1753.E15.gz43_222472
M00056267D:B09	ES 182	562173	1753.G15.gz43_222474
M00056268D:G07	ES 182	639520	1753.H12.gz43_222427
M00056269D:H01	ES 182	639003	1753.I03.gz43_222284

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CloneID	ES No	ClusterID	SequenceName
M00056270A;B09	ES 182	647828	1753.I06.gz43 222332
M00056270B:F02	ES 182	557857	1753.I12.gz43 222428
M00056271A:G04	ES 182	466894	1753.I24.gz43 222620
M00056271C:C06	ES 182	528162	1753,J04.gz43 222301
M00056272D:C03	ES 182	642812	1753.J22.gz43 222589
M00056273A:A07	ES 182	642200	1753.K02.gz43 222270
M00056274D:E08	ES 182	476380	1753.L06.gz43 222335
M00056276A;D05	ES 182	580179	1753.M10.gz43_222400
M00056276A:F05	ES 182	648442	1753.M14.gz43_222464
M00056276D:B12	ES 182	507066	1753.M17.gz43_222512
M00056277B:G05	ES 182	648609	1753,N05.gz43_222321
M00056279B:G06	ES 182	649592	1753.O18.gz43_222530
M00056280B:E09	ES 182	649349	1753.P13.gz43_222451
M00056280D:C06	ES 182	509202	1753.P23.gz43_222611
M00056284C:A11	ES 182	556497	1754.C23.gz43_222982
M00056285B:B01	ES 182	647839	1754.D07.gz43_222727
M00056286A:A12	ES 182	644829	1754.D20.gz43_222935
M00056286B:A04	ES 182	559857	1754.D24.gz43_222999
M00056286C:H08	ES 182	554276	1754.E10.gz43_222776
M00056287C:H08	ES 182	647462	1754.E21.gz43_222952
M00056288A:E12	ES 182	446732	1754.E23.gz43_222984
M00056288D:D03	ES 182	647759	1754.F11.gz43_222793
M00056289B:E05	ES 182	555998	1754.F20.gz43_222937
M00056289B:F09	ES 182	648524	1754.F21.gz43_222953
M00056291B:G01	ES 182	646060	1754.H12.gz43_222811
M00056291C:C11	ES 182	598471	1754.H19.gz43_222923
M00056292D:C06	ES 182	644862	1754.I07.gz43_222732
M00056293B:F02	ES 182	449882	1754.I19.gz43_222924
M00056294B:G06	ES 182	419801	1754.J20.gz43_222941
M00056295C:F12	ES 182	492587	1754.K07.gz43_222734
M00056296A:A07	ES 182	646387	1754.K12.gz43_222814
M00056297D:B10	ES 182	467709	1754.L23.gz43_222991
M00056298A:H02	ES 182	651057	1754.M07.gz43_222736
M00056298B:F10	ES 182	639804	1754.M12.gz43_222816
M00056298D:G01	ES 182	462865	1754.M22.gz43_222976
M00056299C:F06	ES 182	643984	1754.N10.gz43_222785
M00056299D:A02	ES 182	462986	1754.N12.gz43_222817
M00056300A:E06	ES 182	485480	1754.N22.gz43_222977
M00056300B;A11	ES 182	651115	1754.N24.gz43_223009
M00056301C;H08	ES 182	645844	1754.O22.gz43_222978
M00056302D;E12	ES 182	644836	1754.P22.gz43_222979
M00056303A:B03	ES 182	462398	1754.P23.gz43_222995
M00056304A:H04	ES 182	467364	1755.A17.gz43_223268
M00056304B:C05	ES 182	648683	1755.A18.gz43_223284

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00056304C:C03	ES 182	645073	1755.A23.gz43_223364
M00056305A:A02	ES 182	524261	1755.B06.gz43_223093
M00056305B:F09	ES 182	644880	1755.B14.gz43_223221
M00056305D:C05	ES 182	463595	1755.B20.gz43_223317
M00056305D:E08	ES 182	641681	1755.B22.gz43_223349
M00056307A:F07	ES 182	644660	1755.D05.gz43_223079
M00056307D:G02	ES 182	642145	1755.D17.gz43_223271
M00056308A;C01	ES 182	650639	1755.D20.gz43_223319
M00056308C:C12	ES 182	469802	1755.E12.gz43_223192
M00056310A:C10	ES 182	557052	1755.E15.gz43_223240
M00056310B:E06	ES 182	649514	1755.E22.gz43_223352
M00056311C:G06	ES 182	558879	1755.G04.gz43_223066
M00056313A:B06	ES 182	44503	1755.H16.gz43_223259
M00056319A:A11	ES 182	635062	1755,I11.gz43_223180
M00056320C:E08	ES 182	556444	1755.K06.gz43_223102
M00056320D:F07	ES 182	642260	1755.K10.gz43_223166
M00056323A:C09	ES 182	647951	1755.K13.gz43_223214
M00056323A:G03	ES 182	569195	1755.K15.gz43_223246
M00056323A:H05	ES 182	550403	1755.K16.gz43 223262
M00056326C:G07	ES 182	639378	1755.M08.gz43_223136
M00056328B:E07	ES 182	444222	1755.M16.gz43 223264
M00056328D:B01	ES 182	447126	1755.M19.gz43 223312
M00056329C:B09	ES 182	647248	1755.N05.gz43 223089
M00056330B:C02	ES 182	452075	1755.N14.gz43 223233
M00056333A:E09	ES 182	460929	1755.O09.gz43_223154
M00056333C:D08	ES 182	446520	1755.O23.gz43_223378
M00056338B:B08	ES 182	606129	1755.P24.gz43_223395
M00042751B:C07	ES 183	179760	1767.A07.gz43_224579
M00042751D:D03	ES 183	449215	1767.A14.gz43_224691
M00042751D:F09	ES 183	484355	1767.A16.gz43_224723
M00042751D:G06	ES 183	485431	1767.A18.gz43_224755
M00042751D:G09	ES 183	485441	1767.A19.gz43 224771
M00042752A:B04	ES 183	449275	1767.A20.gz43 224787
M00042753B:C08	ES 183	480960	1767.B13.gz43 224676
M00042753C:G12	ES 183	485183	1767.B19.gz43 224772
M00042753D:A12	ES 183	479061	1767.B20.gz43 224788
M00042753D:H11	ES 183	449746	1767.B23.gz43 224836
M00042754B:G02	ES 183	485512	1767.C02.gz43 224501
M00042755A:D09	ES 183	481614	1767.C06.gz43_224565
M00042755A;H04	ES 183	485880	1767.C10.gz43 224629
M00042755B:D01	ES 183	481594	1767.C12.gz43 224661
M00042755B:D09	ES 183	449399	1767.C13.gz43_224677
M00042755B:H04	ES 183	485882	1767.C16.gz43_224725

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00042755B:H11	ES 183	485907	1767.C17.gz43_224741
M00042755C:A10	ES 183	452104	1767.C19.gz43_224773
M00042756B:B01	ES 183	284586	1767.D03.gz43_224518
M00042757A:H07	ES 183	449076	1767.D15.gz43_224710
M00042758B:C06	ES 183	448325	1767.D23.gz43 224838
M00042758B:G04	ES 183	476876	1767.D24.gz43_224854
M00042760A:G12	ES 183	477098	1767.E15.gz43_224711
M00042760B:E12	ES 183	453024	1767.E18.gz43_224759
M00042760C:F10	ES 183	476438	1767.E22.gz43_224823
M00042761B:C11	ES 183	473592	1767.F04.gz43_224536
M00042761B:E05	ES 183	475562	1767.F08.gz43_224600
M00042761C:G11	ES 183	477521	1767.F12.gz43_224664
M00042761D:D12	ES 183	474580	1767.F17.gz43_224744
M00042762A:G02	ES 183	455143	1767.F22.gz43_224824
M00042763B:A03	ES 183	470801	1767.G09.gz43_224617
M00042764A:F12	ES 183	448801	1767.G22.gz43_224825
M00042764B:B10	ES 183	453606	1767.H01.gz43_224490
M00042764B:G10	ES 183	448865	1767.H04.gz43_224538
M00042764C:B10	ES 183	448251	1767.H05.gz43_224554
M00042766A:D07	ES 183	128749	1767.I04.gz43_224539
M00042766B:G11	ES 183	484633	1767.I08.gz43_224603
M00042766D:C05	ES 183	450551	1767.I14.gz43_224699
M00042767A:B10	ES 183	479880	1767.I18.gz43_224763
M00042767B:E10	ES 183	482868	1767.I21.gz43_224811
M00042767B:G04	ES 183	484964	1767,I22.gz43_224827
M00042767B:G09	ES 183	484987	1767,I23.gz43_224843
M00042767D:D02	ES 183	449437	1767.J05.gz43_224556
M00042768A:H09	ES 183	486452	1767,J09.gz43_224620
M00042768B:F02	ES 183	456513	1767.J11.gz43_224652
M00042768C:E06	ES 183	482486	1767.J15.gz43_224716
M00042769B:E12	ES 183	482876	1767.K05.gz43_224557
M00042769C:B01	ES 183	479851	1767.K08.gz43 224605
M00042769C:B06	ES 183	479868	1767.K09.gz43_224621
M00042770A:D02	ES 183	449438	1767.K15.gz43 224717
M00042771B:G05	ES 183	485073	1767.L12.gz43 224670
M00042771D:F02	ES 183	449592	1767.L16.gz43 224734
M00042772C:C12	ES 183	447218	1767.L20.gz43_224798
M00042773A:G07	ES 183	485086	1767.M02.gz43 224511
M00042773A;H11	ES 183	449689	1767.M03.gz43 224527
M00042773B;H08	ES 183	486547	1767.M06.gz43 224575
M00042774B:C01	ES 183	481057	1767.M14.gz43_224703
M00042774C;B09	ES 183	449261	1767.M16.gz43 224735
M00042775B:C09	ES 183	480703	1767.N04.gz43_224544
M00042776D:D01	ES 183	448152	1767.N17.gz43 224752

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00042776D:G10	ES 183	477399	1767.N18.gz43_224768
M00042777A:B10	ES 183	472119	1767.N20.gz43_224800
M00042777B:B05	ES 183	472101	1767.O02.gz43_224513
M00042777B:H02	ES 183	449015	1767.O03.gz43_224529
M00042778A:C04	ES 183	473433	1767.O11.gz43_224657
M00042778A:G07	ES 183	477387	1767.O12.gz43_224673
M00042778B:B07	ES 183	448260	1767.O15.gz43_224721
M00042778D:F11	ES 183	475797	1767.O23.gz43_224849
M00042779A;B04	ES 183	472425	1767.O24.gz43_224865
M00042779D:D04	ES 183	453385	1767.P15.gz43_224722
M00042779D:E06	ES 183	474823	1767.P17.gz43_224754
M00042780A:H05	ES 183	477757	1767.P24.gz43_224866
M00042809D:C12	ES 183	649735	1777.A23.gz43_252199
M00042809D:G09	ES 183	557375	1777,B01.gz43 251848
M00042811A:A01	ES 183	500896	1777.B23.gz43 252200
M00042811C:B06	ES 183	537506	1777.C03.gz43 251881
M00042812B:C01	ES 183	456793	1777.C11.gz43_252009
M00042815C:C02	ES 183	448340	1777.E11.gz43 252011
M00042815C:F10	ES 183	448793	1777.E13.gz43 252043
M00042816A:C09	ES 183	735534	1777.E17.gz43 252107
M00042816A:G04	ES 183	639163	1777.E21.gz43 252171
M00042816B:C08	ES 183	545162	1777.F03.gz43 251884
M00042817C:B04	ES 183	727517	1777.G05.gz43 251917
M00042819C:H02	ES 183	449718	1777.H14.gz43 252062
M00042821C:H06	ES 183	485653	1777.I17.gz43 252111
M00042824A:C02	ES 183	52034	1777.K05.gz43 251921
M00042825A:B05	ES 183	449258	1777.K17.gz43 252113
M00042828D:B11	ES 183	472400	1777.M09.gz43 251987
M00042831A:F12	ES 183	484207	1777.N18.gz43_252132
M00042831D:G06	ES 183	484669	1777.O01.gz43 251861
M00042833C:G05	ES 183	485020	1777.O18.gz43 252133
M00042835D:C04	ES 183	450883	1777.P19.gz43_252150
M00042836A:B12	ES 183	480227	1777.P24.gz43 252230
M00042836D:C07	ES 183	449291	1778.A11.gz43 225411
M00042838A:E02	ES 183	475111	1778.B02.gz43 225268
M00042839B:F05	ES 183	475942	1778.B23.gz43 225604
M00042839D:F02	ES 183	476256	1778.C07.gz43 225349
M00042840B:F08	ES 183	452202	1778.C12.gz43 225429
M00042840C:B12	ES 183	447517	1778.C14.gz43 225461
M00042841B:H05	ES 183	450673	1778.C18.gz43 225525
M00042841D.G10	ES 183	446657	1778,C23,gz43 225605
M00042841B:G10	ES 183	472307	1778.C24.gz43 225621
M00042842A:B12	ES 183	448233	1778.D01.gz43_225254
M00042842C:H11	ES 183	477555	1778.D12.gz43_225430

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CloneID	ES No	ClusterID	SequenceName
M00042843A:E07	ES 183	474597	1778.D18.gz43_225526
M00042843B;F05	ES 183	453856	1778.D22.gz43_225590
M00042844A:D02	ES 183	447983	1778.E07.gz43_225351
M00042844C:B03	ES 183	448230	1778.E12.gz43_225431
M00042844C:C12	ES 183	448383	1778.E14.gz43_225463
M00042844D:H02	ES 183	477593	1778.E18.gz43_225527
M00042845A:B05	ES 183	448104	1778.E19.gz43_225543
M00042845C:A09	ES 183	447983	1778.F03.gz43_225288
M00042846C:D09	ES 183	639991	1778.F23.gz43_225608
M00042847A:F04	ES 183	727875	1778.G10.gz43_225401
M00042848D:G12	ES 183	448029	1778.H06.gz43_225338
M00042849B:G06	ES 183	485029	1778.H12.gz43_225434
M00042850A:B11	ES 183	404603	1778.H20.gz43_225562
M00042850C:C10	ES 183	646583	1778,I06.gz43_225339
M00042851A:B08	ES 183	449273	1778.I11.gz43_225419
M00042851A:E11	ES 183	483274	1778.I12.gz43_225435
M00042851D:B08	ES 183	452990	1778.I18.gz43_225531
M00042852B:C06	ES 183	480900	1778.J04.gz43_225308
M00042852D:G07	ES 183	485480	1778.J10.gz43_225404
M00042853A:G03	ES 183	644190	1778.J12.gz43_225436
M00042854A:B11	ES 183	551811	1778.K04.gz43_225309
M00042854A:D05	ES 183	725641	1778.K07.gz43_225357
M00042854B:F05	ES 183	725266	1778.K12.gz43_225437
M00042854D:A05	ES 183	482071	1778.K16.gz43_225501
M00042855B:H06	ES 183	485841	1778.L05.gz43_225326
M00042855C:G11	ES 183	484789	1778.L08.gz43_225374
M00042855D:A12	ES 183	449110	1778.L09.gz43_225390
M00042855D:B12	ES 183	480005	1778.L12.gz43_225438
M00042855D:D06	ES 183	456737	1778.L14.gz43_225470
M00042857C:G04	ES 183	449652	1778.M11.gz43_225423
M00042857C:G05	ES 183	484841	1778.M12.gz43_225439
M00042858A:A08	ES 183	479084	1778.M14.gz43_225471
M00042859B:D06	ES 183	449403	1778.N02.gz43_225280
M00042859D:D12	ES 183	481664	1778.N11.gz43_225424
M00042860A:C01	ES 183	480641	1778.N12.gz43_225440
M00042860A:H04	ES 183	485924	1778.N16.gz43_225504
M00042860C:E02	ES 183	483066	1778.N23,gz43_225616
M00042860D:F02	ES 183	449604	1778.O02.gz43_225281
M00042860D:F05	ES 183	484091	1778.003.gz43_225297
M00042860D:H10	ES 183	449751	1778,O06,gz43_225345
M00042861A:F04	ES 183	451368	1778.O09.gz43_225393
M00042861C:C03	ES 183	481319	1778.O13.gz43_225457
M00042861C:F07	ES 183	484468	1778,O15.gz43_225489
M00042861D:D09	ES 183	449407	1778.O18.gz43_225537

Table 13

CloneID M00042862A:C01 M00042862C:E02	ES No ES 183	ClusterID	SequenceName
M00042862C:E02	ES 183		
		480640	1778.O23.gz43_225617
	ES 183	449521	1778.P09.gz43 225394
M00056352D:B04	ES 183	559486	1780.A05.gz43_226083
M00056352D:H02	ES 183	594994	1780.A10.gz43_226163
M00056353C:E12	ES 183	736210	1780.B04.gz43_226068
M00056353D:D10	ES 183	731925	1780.B11.gz43_226180
M00056355A:C10	ES 183	463966	1780.C10.gz43_226165
M00056355B:D02	ES 183	557029	1780.C16.gz43_226261
M00056355C:A05	ES 183	553951	1780.C18.gz43_226293
M00056355D:A05	ES 183	727129	1780.D07.gz43_226118
M00056355D:B08	ES 183	468467	1780.D09.gz43_226150
M00056355D:G04	ES 183	727235	1780.D11.gz43_226182
M00056356C:H07	ES 183	732150	1780.E02.gz43_226039
M00056357A:C08	ES 183	551167	1780.E10.gz43_226167
M00056357C:C06	ES 183	724039	1780.E23.gz43_226375
M00056358A:A10	ES 183	723859	1780.F10.gz43_226168
M00056358A:G05	ES 183	734150	1780.F12.gz43_226200
M00056358D:A06	ES 183	465697	1780.F22.gz43_226360
M00056359A:C08	ES 183	623978	1780.G05.gz43_226089
M00056359A:F06	ES 183	416326	1780.G06.gz43_226105
M00056360A:D11	ES 183	561719	1780.H08.gz43_226138
M00056360B:D09	ES 183	528981	1780.H12.gz43_226202
M00056360C:A03	ES 183	723869	1780.H14.gz43_226234
M00056360C:B01	ES 183	480005	1780.H15.gz43_226250
M00056360D:H10	ES 183	732144	1780.H23.gz43_226378
M00056361A:C01	ES 183	733806	1780.I01.gz43_226027
M00056361A:H06	ES 183	600856	1780.I05.gz43_226091
M00056361B:E02	ES 183	471646	1780.I10.gz43_226171
M00056361C:C06	ES 183	135364	1780.I12.gz43_226203
M00056362C:C02	ES 183	511606	1780.J12.gz43_226204
M00056363A:C06	ES 183	724907	1780.K02.gz43_226045
M00056363A:E10	ES 183	364334	1780.K05.gz43_226093
M00056363C:C12	ES 183	552891	1780.K14.gz43_226237
M00056363D:H03	ES 183	728774	1780.K22.gz43_226365
M00056364A:B04	ES 183	724792	1780,L02.gz43_226046
M00056364C:B01	ES 183	503862	1780.L08.gz43_226142
M00056364C:H08	ES 183	588959	1780,L15.gz43_226254
M00056364D:C05	ES 183	553244	1780.L17.gz43_226286
M00056365B:G10	ES 183	480924	1780.M11.gz43_226191
M00056366B:A11	ES 183	449242	1780.N05.gz43_226096
M00056367C:D06	ES 183	560859	1780.007.gz43_226129
M00056367C:H05	ES 183	642558	1780.O12.gz43_226209
M00056368A:C11	ES 183	456535	1780.O20.gz43_226337
M00056368B:H09	ES 183	661194	1780.P05.gz43_226098

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ES No	ClusterID	SequenceName
	482831	1780.P07.gz43_226130
ES 184	517298	1780.P11.gz43_226194
ES 184	730301	1780.P14.gz43_226242
ES 184	728389	1780.P22.gz43_226370
ES 184	736001	1789.A03.gz43_226435
ES 184	734646	1789.B02.gz43_226420
ES 184	568484	1789.B04.gz43_226452
ES 184	503625	1789.B08.gz43_226516
ES 184	735649	1789.B11.gz43_226564
ES 184	734943	1789.C12.gz43_226581
ES 184	725389	1789.D04.gz43_226454
ES 184	725381	1789.E09.gz43_226535
ES 184	487079	1789.F03.gz43_226440
ES 184	734348	1789.F11.gz43_226568
ES 184	724257	1789.G01.gz43_226409
ES 184	649411	1789.G09.gz43_226537
ES 184	460284	1789.H03.gz43 226442
ES 184	724694	1789.H12.gz43 226586
ES 184	725274	1789.I03.gz43_226443
ES 184	729993	1789.I09.gz43 226539
ES 184	723972	1789.J12.gz43 226588
ES 184	503491	1789.K03.gz43 226445
ES 184	731542	1789.K05.gz43 226477
ES 184	725321	1789.L04.gz43_226462
ES 184	725448	1789.L06.gz43 226494
ES 184	549790	1789.L07.gz43 226510
ES 184	474346	1789.M01.gz43_226415
ES 184	186372	1789.M02.gz43_226431
ES 184	727523	1789.M10.gz43_226559
ES 184	726636	1789.N01.gz43 226416
ES 184	724388	1789.N02.gz43 226432
ES 184	553035	1789.N03.gz43 226448
ES 184	452806	1789.N06.gz43 226496
ES 184	730660	1789;N07.gz43 226512
ES 184	459918	1789.N10.gz43_226560
ES 184	623408	1789.N12.gz43 226592
ES 184	558412	1789,O11.gz43 226577
ES 184	475682	1789.P01.gz43 226418
ES 184	724009	1789.P03.gz43 226450
ES 184	729637	1790.A10.gz43 226931
ES 184	495712	1790,A17.gz43 227043
ES 184	725165	1790,B08.gz43 226900
ES 184	556769	1790.B11.gz43_226948
	ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184 ES 184	ES 184 482831 ES 184 517298 ES 184 730301 ES 184 730301 ES 184 736001 ES 184 736001 ES 184 736001 ES 184 736001 ES 184 736001 ES 184 736001 ES 184 735649 ES 184 735649 ES 184 725381 ES 184 725381 ES 184 725381 ES 184 734948 ES 184 734948 ES 184 734948 ES 184 734948 ES 184 734948 ES 184 734948 ES 184 734948 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349694 ES 184 7349696 ES 184 7349696 ES 184 7349696 ES 184 736660 ES 184 736660 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690 ES 184 736690

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CloneID	ES No	ClusterID	SequenceName
M00056390C:C04	ES 184	724914	1790.B12.gz43_226964
M00056391C:H12	ES 184	727150	1790.C14.gz43_226997
M00056391D:E06	ES 184	457508	1790.C18.gz43_227061
M00056392A;D11	ES 184	477296	1790.C19.gz43_227077
M00056392A:F06	ES 184	448817	1790,C20.gz43_227093
M00056392D:C05	ES 184	426297	1790.D07.gz43_226886
M00056392D:H02	ES 184	729111	1790.D08.gz43_226902
M00056393A:F01	ES 184	549480	1790.D11.gz43_226950
M00056393A:G06	ES 184	614463	1790.D12.gz43_226966
M00056393D:A09	ES 184	558794	1790.D21.gz43_227110
M00056393D:C12	ES 184	118514	1790.D24.gz43_227158
M00056394B:C11	ES 184	503173	1790.E14.gz43_226999
M00056394B:G05	ES 184	725454	1790.E19.gz43_227079
M00056394C:D07	ES 184	640158	1790.F01.gz43_226792
M00056395C:H03	ES 184	523364	1790.F23.gz43_227144
M00056395D:B01	ES 184	732821	1790.F24.gz43_227160
M00056396A:D03	ES 184	733563	1790.G08.gz43_226905
M00056396A:G10	ES 184	474981	1790.G12.gz43_226969
M00056396B:E10	ES 184	642985	1790.G16.gz43_227033
M00056396D:H01	ES 184	484934	1790.H05.gz43_226858
M00056397C:A09	ES 184	515038	1790.H17.gz43_227050
M00056398A:B10	ES 184	724851	1790.I11.gz43_226955
M00056398C:B09	ES 184	735396	1790.I24.gz43_227163
M00056398C:D10	ES 184	725095	1790.J02.gz43_226812
M00056399B:G09	ES 184	448758	1790,J15.gz43_227020
M00056399D:C11	ES 184	656268	1790.K04.gz43_226845
M00056400A:D08	ES 184	726494	1790.K10.gz43_226941
M00056400A:G10	ES 184	725451	1790.K16.gz43_227037
M00056400C:B11	ES 184	447222	1790.K24.gz43_227165
M00056400C:G04	ES 184	479851	1790,L03,gz43_226830
M00056401A:F09	ES 184	729950	1790.L12.gz43_226974
M00056401B:B01	ES 184	735234	1790.L15.gz43_227022
M00056402B:D07	ES 184	641576	1790.M06.gz43_226879
M00056402C:B09	ES 184	649591	1790.M07.gz43_226895
M00056402C:E10	ES 184	483008	1790.M09.gz43_226927
M00056402D:A01	ES 184	724714	1790.M10.gz43_226943
M00056402D:B07	ES 184	197839	1790.M11.gz43_226959
M00056403A;E09	ES 184	417549	1790.M18.gz43_227071
M00056403B:D09	ES 184	468467	1790.M23.gz43_227151
M00056403B:G12	ES 184	729779	1790.M24.gz43_227167
M00056403C:F02	ES 184	481077	1790.N03.gz43_226832
M00056403C;F03	ES 184	724411	1790.N04.gz43_226848
M00056403C:F12	ES 184	640261	1790.N05.gz43_226864
M00056404A:G03	ES 184	640756	1790.N12.gz43_226976

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1 able 13			
CloneID	ES No	ClusterID	SequenceName
M00056404B:B06	ES 184	188309	1790,N14.gz43_227008
M00056404C;C12	ES 184	733643	1790.N22.gz43_227136
M00056404D:G03	ES 184	562769	1790.O02.gz43_226817
M00056405C:C12	ES 184	724078	1790.O11.gz43_226961
M00056405C:H04	ES 184	449228	1790.O13.gz43_226993
M00056405D:E05	ES 184	730948	1790.O17.gz43_227057
M00056406A:C02	ES 184	726411	1790.O20.gz43 227105
M00056406A;E01	ES 184	728454	1790,O23.gz43_227153
M00056406B:C05	ES 184	456577	1790.P03.gz43_226834
M00056406C:D09	ES 184	543323	1790.P10.gz43_226946
M00056406C:F12	ES 184	423420	1790.P13.gz43_226994
M00056407A:D10	ES 184	483147	1790.P15.gz43_227026
M00056407A:G03	ES 184	729809	1790.P16.gz43_227042
M00056409A:C03	ES 184	641687	1790.P21.gz43_227122
M00056409A;D03	ES 184	470462	1790.P23.gz43_227154
M00056409A:D06	ES 184	727633	1790.P24.gz43_227170
M00056409B:C12	ES 184	729784	1791.A03.gz43_227203
M00056409B:E07	ES 184	734253	1791.A05.gz43_227235
M00056409C:B04	ES 184	711493	1791.A06.gz43_227251
M00056410A:A04	ES 184	640744	1791.A19.gz43_227459
M00056410B:A09	ES 184	641383	1791.A23.gz43_227523
M00056410D:H03	ES 184	49703	1791.B14.gz43_227380
M00056411A:H06	ES 184	558049	1791.B17.gz43_227428
M00056411B;D01	ES 184	703978	1791.B19.gz43_227460
M00056412B:B08	ES 184	724773	1791.C22.gz43_227509
M00056413B:D07	ES 184	642637	1791.D18.gz43_227446
M00056413C:B06	ES 184	723951	1791.D24.gz43 227542
M00056414C:H08	ES 184	729899	1791,E18.gz43_227447
M00056415A:D03	ES 184	475797	1791.F01.gz43_227176
M00056415B:B10	ES 184	402471	1791.F03.gz43_227208
M00056416B:B11	ES 184	648170	1791.G01.gz43_227177
M00056416B:C09	ES 184	456544	1791.G02.gz43_227193
M00056416B:H01	ES 184	727090	1791.G04.gz43_227225
M00056416D:C08	ES 184	729199	1791.G12.gz43_227353
M00056416D:D12	ES 184	435349	1791.G13.gz43 227369
M00056416D:H10	ES 184	728986	1791.G18.gz43 227449
M00056417A:E08	ES 184	733484	1791.G23.gz43_227529
M00056417B:D05	ES 184	642170	1791.H05.gz43_227242
M00056417C:G10	ES 184	725408	1791.H14.gz43_227386
M00056417D:G09	ES 184	725407	1791.H20.gz43_227482
M00056418A:G01	ES 184	729579	1791.H24.gz43_227546
M00056418A:G09	ES 184	472684	1791.I01.gz43_227179
M00056419A:F08	ES 184	542957	1791.J07.gz43_227276
M00056419D:D02	ES 184	728440	1791.J24.gz43_227548

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CloneID	ES No	ClusterID	SequenceName
M00056419D:G04	ES 184	453762	1791.K03.gz43_227213
M00056420A:C01	ES 184	560957	1791.K04.gz43_227229
M00056420A:C12	ES 184	128749	1791.K05.gz43_227245
M00056420A;E09	ES 184	643646	1791.K07.gz43_227277
M00056420A:G11	ES 184	731390	1791.K10.gz43_227325
M00056420B:G05	ES 184	480377	1791.K12.gz43_227357
M00056420C:E08	ES 184	736507	1791.K21.gz43_227501
M00056420D:A09	ES 184	562581	1791.K23.gz43_227533
M00056421A:F12	ES 184	646146	1791.L05.gz43_227246
M00056421C:E06	ES 184	562137	1791.L11.gz43_227342
M00056421C:E12	ES 184	730178	1791.L12.gz43_227358
M00056421C:H01	ES 184	500253	1791.L14.gz43_227390
M00056422A:C12	ES 184	737019	1791.L22.gz43_227518
M00056422A:F07	ES 184	498194	1791.L24.gz43_227550
M00056422B:A08	ES 184	511149	1791.M03.gz43_227215
M00056422B:C12	ES 184	727845	1791.M04.gz43_227231
M00056422C:B10	ES 184	474965	1791.M15.gz43_227407
M00056422D:D10	ES 184	732987	1791.M22.gz43_227519
M00056423B:A12	ES 184	650437	1791.N06.gz43_227264
M00056423C:G05	ES 184	644723	1791.N14.gz43_227392
M00056423C:H03	ES 184	651053	1791.N15.gz43_227408
M00056423D:B06	ES 184	724024	1791.N17.gz43_227440
M00056423D:F10	ES 184	288134	1791.N24.gz43_227552
M00056423D:H07	ES 184	477718	1791.O02.gz43_227201
M00056424A:F05	ES 184	733490	1791.O08.gz43_227297
M00056424A:F12	ES 184	649722	1791.O09.gz43_227313
M00056424B:A11	ES 184	736893	1791.O13.gz43_227377
M00056424C:B10	ES 184	725408	1791.O19.gz43_227473
M00056424C:F02	ES 184	555359	1791.O21.gz43_227505
M00056424C:H04	ES 184	437450	1791.O24.gz43_227553
M00056424D:C08	ES 184	736632	1791.P05.gz43_227250
M00056424D:C12	ES 184	674574	1791.P06.gz43_227266
M00056425B:B02	ES 184	723963	1791.P13.gz43_227378
M00056425D:D08	ES 184	724220	1791.P22.gz43_227522
M00056425D:H01	ES 184	726768	1791.P23.gz43_227538
M00056426B:G12	ES 184	731158	1792.A09.gz43_227695
M00056426C:G05	ES 184	452981	1792.A11.gz43_227727
M00056427C:E06	ES 184	732827	1792.A21.gz43_227887
M00056428A:F10	ES 184	592186	1792.B09.gz43_227696
M00056428B:B01	ES 184	710362	1792.B11.gz43_227728
M00056429D:E02	ES 184	480005	1792.C11.gz43_227729
M00056432A;A01	ES 184	556115	1792.D06.gz43_227650
M00056432A:C08	ES 184	472073	1792.D08.gz43_227682
M00056433B:A09	ES 184	724400	1792.E03.gz43_227603

Table 13

Table 13			
CloneID	ES No	ClusterID	SequenceName
M00056433D:A09	ES 184	554398	1792.E12.gz43_227747
M00056433D:D08	ES 184	570928	1792.E14.gz43_227779
M00056434C:B01	ES 184	734287	1792.F06.gz43_227652
M00056435B:E06	ES 184	455972	1792.F20.gz43_227876
M00056435C:D04	ES 184	492089	1792.F22.gz43_227908
M00056476C:E05	ES 184	725238	1792.G10.gz43_227717
M00056476D:H11	ES 184	647375	1792.G16.gz43_227813
M00056477A:B11	ES 184	736751	1792.G17.gz43_227829
M00056477C:G09	ES 184	725454	1792.H12.gz43_227750
M00056478C:H01	ES 184	725584	1792.I07.gz43_227671
M00056478D:G02	ES 184	729584	1792.I12.gz43_227751
M00056479C:E04	ES 184	645476	1792.J05.gz43_227640
M00056480B:D06	ES 184	482461	1792.K02.gz43_227593
M00056481A:F02	ES 184	734646	1792.K19.gz43_227865
M00056482B:B11	ES 184	481594	1792.L18.gz43_227850
M00056482B:C09	ES 184	453713	1792.L20.gz43_227882
M00056482B:H02	ES 184	466795	1792.L22.gz43_227914
M00056483A:F10	ES 185	452618	1792.M09.gz43_227707
M00056483B:E04	ES 185	724183	1792.M13.gz43_227771
M00056483B:F04	ES 185	724390	1792.M14.gz43_227787
M00056483C:D06	ES 185	724183	1792.M20.gz43_227883
M00056483D:F06	ES 185	730282	1792.M24.gz43 227947
M00056484B:E11	ES 185	725210	1792.N12.gz43 227756
M00056484D:D01	ES 185	675299	1792.N18.gz43_227852
M00056484D:G08	ES 185	650940	1792.N21.gz43 227900
M00056485B:D11	ES 185	451404	1792.004.gz43 227629
M00056486A:C12	ES 185	477296	1792.O16.gz43 227821
M00056486B:G02	ES 185	553983	1792.P01.gz43 227582
M00056486C:F04	ES 185	717743	1792.P05.gz43 227646
M00056486D:A12	ES 185	725006	1792,P07.gz43 227678
M00056486D:F04	ES 185	727948	1792.P10.gz43 227726
M00056487A:C10	ES 185	471931	1792.P11.gz43 227742
M00056487A:D01	ES 185	462687	1792.P12.gz43 227758
M00056487A;F05	ES 185	725348	1792.P17.gz43 227838
M00056487C:E03	ES 185	731820	1801.A04.gz43 227999
M00056487C:E04	ES 185	449291	1801.A05.gz43 228015
M00056488B:E04	ES 185	462266	1801.B03.gz43 227984
M00056488B:G10	ES 185	725434	1801.B05.gz43 228016
M00056488B:H04	ES 185	429191	1801.B06.gz43 228032
M00056491B:B09	ES 185	733006	1801.D01.gz43 227954
M00056491B:E02	ES 185	447785	1801.D02.gz43 227970
M00056491B:G05	ES 185	472485	1801.D04.gz43 228002
M00056491B:G08	ES 185	446191	1801.D05.gz43 228018
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Table 13

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M00056492B:D06	ES 185	446928	1801.E04.gz43_228003
M00056495A:A07	ES 185	447658	1801.G03.gz43_227989
M00056495A:C02	ES 185	460690	1801.G05.gz43_228021
M00056496D:B12	ES 185	733151	1801.I01.gz43_227959
M00056496D:G03	ES 185	649202	1801.I05.gz43_228023
M00056497C:D05	ES 185	734261	1801.J01.gz43_227960
M00056497C:E01	ES 185	523753	1801.J02.gz43_227976
M00056497D:C11	ES 185	611604	1801.J05.gz43_228024
M00056498D:C01	ES 185	514142	1801.K01.gz43_227961
M00056499A:A04	ES 185	635951	1801.K03.gz43_227993
M00056499A:B10	ES 185	447002	1801.K05.gz43_228025
M00056500A:G12	ES 185	635951	1801.L05.gz43_228026
M00056501B:B09	ES 185	471232	1801.M01.gz43_227963
M00056501B:C07	ES 185	451383	1801.M02.gz43_227979
M00056503B:G11	ES 185	640116	1801.006.gz43_228045
M00056504B:B01	ES 185	732598	1801.P04.gz43_228014
M00056504B:C03	ES 185	736665	1801.P05.gz43_228030
M00056567A:H06	ES 185	727410	1813.A13.gz43_229679
M00056568C:D02	ES 185	454560	1813.B14.gz43_229696
M00056570A:D09	ES 185	548893	1813.C22.gz43_229825
M00056572A:A04	ES 185	736194	1813.E06.gz43_229571
M00056572C:E03	ES 185	458186	1813.E22.gz43_229827
M00056573C:B09	ES 185	554557	1813.F20.gz43_229796
M00056576A:A04	ES 185	734043	1813.H05.gz43_229558
M00056576A:E01	ES 185	475757	1813.H08.gz43_229606
M00056576C:G01	ES 185	513540	1813,H22.gz43_229830
M00056577A:F10	ES 185	508210	1813.I06.gz43_229575
M00056577D:F08	ES 185	733664	1813.I20.gz43_229799
M00056578A:B05	ES 185	725759	1813.I22.gz43_229831
M00056578D:A02	ES 185	729281	1813.J10.gz43_229640
M00056581D:A08	ES 185	465446	1813.L13.gz43_229690
M00056583C:A05	ES 185	511351	1813.M13.gz43_229691
M00056584C:A06	ES 185	733970	1813.N10.gz43_229644
M00056585C:C08	ES 185	727260	1813.007.gz43_229597
M00056586B:D10	ES 185	461062	1813.O24.gz43_229869
M00056631D:F06	ES 185	730897	1816,A24.gz43_231007
M00056632C:E03	ES 185	736325	1816.B20.gz43_230944
M00056632C:H08	ES 185	726494	1816.B22.gz43_230976
M00056632D:E02	ES 185	727944	1816,B24.gz43_231008
M00056633D:E05	ES 185	726104	1816.C19.gz43_230929
M00056634A:C06	ES 185	736778	1816.C21.gz43_230961
M00056634A:G11	ES 185	726448	1816.C23.gz43_230993
M00056636C:H05	ES 185	482868	1816.E19.gz43_230931
M00056636D:F03	ES 185	559684	1816.E21.gz43_230963

Table 13

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ClonelD	ES No	ClusterID	SequenceName
M00056637A:C08	ES 185	647808	1816.E24.gz43_231011
M00056639A;E07	ES 185	500233	1816.G20.gz43_230949
M00056639A:F11	ES 185	419711	1816.G22.gz43_230981
M00056639A:H12	ES 185	215249	1816.G23.gz43_230997
M00056640C;C01	ES 185	196279	1816.H21.gz43_230966
M00056641C:G09	ES 185	727332	1816.I19.gz43_230935
M00056642D:C05	ES 185	726818	1816.J20.gz43_230952
M00056644C:B03	ES 185	727078	1816,K19,gz43_230937
M00056645D:E06	ES 185	727108	1816.L19.gz43_230938
M00056645D:E08	ES 185	727110	1816.L20.gz43_230954
M00056645D:G01	ES 185	659700	1816,L22,gz43_230986
M00056646A:B01	ES 185	733375	1816.L24.gz43_231018
M00056646D:E03	ES 185	724083	1816.M20.gz43_230955
M00056646D:F07	ES 185	735392	1816.M22.gz43_230987
M00056647A:F06	ES 185	730596	1816.M24.gz43_231019
M00056648A:H05	ES 185	481930	1816.N19.gz43_230940
M00056648B:C08	ES 185	724565	1816.N20.gz43_230956
M00056648B:E09	ES 185	735322	1816.N23.gz43_231004
M00056650A:E01	ES 185	207530	1816.O20.gz43_230957
M00056650B;C11	ES 185	727175	1816.023.gz43_231005
M00056651B:E12	ES 185	731966	1816.P23.gz43_231006
M00056652B:E09	ES 185	728936	1825.A14.gz43_231231
M00056653D:E07	ES 185	462409	1825,B19,gz43_231312
M00056653D:F07	ES 185	730570	1825,B21.gz43_231344
M00056654A:A07	ES 185	736415	1825.B24.gz43_231392
M00056656A:E01	ES 185	727108	1825.D21.gz43_231346
M00056657A:F06	ES 185	477718	1825.E16.gz43_231267
M00056657C:B04	ES 185	554713	1825.E20.gz43_231331
M00056658A:E11	ES 185	484987	1825.F14.gz43_231236
M00056658B;C03	ES 185	552927	1825.F16.gz43_231268
M00056658C:B12	ES 185	463741	1825.F22.gz43_231364
M00056660B:B06	ES 185	646504	1825.H19.gz43_231318
M00056660B:G12	ES 185	562984	1825.H21.gz43_231350
M00056660C:A06	ES 185	729732	1825.H22.gz43_231366
M00056661D:D06	ES 185	447074	1825.I21.gz43_231351
M00056663A:B02	ES 185	719620	1825.J16.gz43_231272
M00056663A:H09	ES 185	729675	1825.J20.gz43_231336
M00056664A;F09	ES 185	537451	1825.K17.gz43_231289
M00056664D:H06	ES 185	448104	1825.L19.gz43_231322
M00056665A;G08	ES 185	727343	1825.L23.gz43_231386
M00056666A:C04	ES 185	481592	1825.M14.gz43_231243
M00056666A;D09	ES 185	726953	1825.M16.gz43_231275
M00056666A:D10	ES 185	650973	1825.M17.gz43_231291
M00056667A:A12	ES 185	725607	1825.N13.gz43_231228

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Table 13

CloneID	ES No	ClusterID	SequenceName
M00056667B;E04	ES 185	725381	1825.N17.gz43 231292
M00056667C:H01	ES 185	476477	1825.N21.gz43 231356
M00056667D:E05	ES 185	732351	1825.N22.gz43 231372
M00056668D:E03	ES 185	735605	1825.O14.gz43 231245
M00056669B:E10	ES 185	554597	1825.024.gz43 231405
M00056670A:G02	ES 185	45934	1825.P13.gz43 231230
M00056670B;A12	ES 185	734184	1825.P14.gz43 231246
M00056670B;G05	ES 185	478458	1825.P17.gz43 231294
M00056672A:E05	ES 185	723914	1826.A13.gz43_231599
M00056672D:B08	ES 185	726722	1826.A21.gz43 231727
M00056672D:E04	ES 185	546121	1826.A23.gz43 231759
M00056674D:E09	ES 185	734622	1826.C14,gz43 231617
M00056675A;C02	ES 185	446575	1826.C18.gz43 231681
M00056675A;G02	ES 185	513156	1826.C23.gz43 231761
M00056676B:G12	ES 185	641777	1826.D16.gz43 231650
M00056676C:B04	ES 185	725899	1826.D18.gz43_231682
M00056678A:B02	ES 185	482722	1826.E13.gz43_231603
M00056678A:H05	ES 185	43352	1826.E16.gz43 231651
M00056678B:E11	ES 185	734828	1826.E17.gz43 231667
M00056678B:H02	ES 185	736246	1826.E20.gz43 231715
M00056678B:H06	ES 185	725266	1826.E21.gz43 231731
M00056678C:A12	ES 185	730664	1826.E22.gz43 231747
M00056679C:F11	ES 185	736194	1826.F22.gz43 231748
M00056681A:E06	ES 185	729387	1826.G22.gz43 231749
M00056681B;A11	ES 185	728791	1826.G24.gz43 231781
M00056682A:F05	ES 185	447489	1826,H17,gz43 231670
M00056683B:H04	ES 185	645925	1826,I14,gz43 231623
M00056683C:H10	ES 185	644445	1826.I17.gz43_231671
M00056683C:H11	ES 185	728659	1826.I18.gz43_231687
M00056684B:B12	ES 185	559610	1826.I24.gz43 231783
M00056685A:H11	ES 185	517274	1826.J17.gz43 231672
M00056685B:G04	ES 185	473238	1826,J20,gz43 231720
M00056685C;G07	ES 185	729205	1826,J24,gz43 231784
M00056686D:D07	ES 185	733789	1826,K18,gz43 231689
M00056686D:E04	ES 185	730829	1826,K20,gz43 231721
M00056689B;F03	ES 185	492094	1826.M19.gz43_231707
M00056689C:F05	ES 185	729413	1826.M24.gz43 231787
M00056692A;A05	ES 185	492629	1826.O21.gz43 231741
M00056693B:D07	ES 185	727965	1827,A02,gz43_231807
M00056693B:D11	ES 185	580601	1827.A04.gz43_231839
M00056693B:H03	ES 185	573733	1827.A09.gz43_231919
M00056693C:D12	ES 185	727968	1827.A13.gz43_231983
M00056693C:G06	ES 185	617813	1827.A14.gz43_231999
M00056693D:B02	ES 185	425396	1827.A15.gz43_232015

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00056693D:E12	ES 185	475942	1827.A18.gz43_232063
M00056694A:C05	ES 185	553483	1827.A22.gz43_232127
M00056694A:G02	ES 185	479997	1827.B02.gz43_231808
M00056694A:G03	ES 185	693869	1827.B03.gz43_231824
M00056694A:G11	ES 185	493830	1827.B04.gz43_231840
M00056694B:H10	ES 185	730375	1827.B08.gz43_231904
M00056695A:H09	ES 185	449701	1827.B21.gz43_232112
M00056695C:C04	ES 185	561301	1827.C01.gz43_231793
M00056695C:H08	ES 185	540618	1827.C06.gz43_231873
M00056695D:E05	ES 185	472837	1827.C10.gz43_231937
M00056695D:F01	ES 185	735974	1827.C11.gz43_231953
M00056696A:G12	ES 185	650542	1827.C16.gz43_232033
M00056696B:B03	ES 185	727506	1827.C17.gz43_232049
M00056696B:H01	ES 185	551096	1827.C24.gz43_232161
M00056697A:D02	ES 185	588059	1827.D11.gz43_231954
M00056697A:G03	ES 185	395596	1827.D13.gz43 231986
M00056697A:H10	ES 185	725084	1827.D16.gz43 232034
M00056697C:D11	ES 185	727845	1827.D21.gz43_232114
M00056697C:H10	ES 185	728556	1827.E01.gz43_231795
M00056697D:B09	ES 185	724714	1827.E04.gz43_231843
M00056697D:C09	ES 185	501030	1827.E06.gz43 231875
M00056697D:C12	ES 185	462687	1827.E07.gz43 231891
M00056698A:A07	ES 185	530656	1827.E10.gz43 231939
M00056698B:E10	ES 185	540618	1827.E17.gz43 232051
M00056698D:E11	ES 185	737109	1827.F04.gz43 231844
M00056698D:G09	ES 185	477366	1827.F06.gz43_231876
M00056699A:C07	ES 185	454844	1827.F10.gz43_231940
M00056699A;D08	ES 185	452662	1827.F11.gz43_231956
M00056699A:E08	ES 185	672032	1827.F12.gz43_231972
M00056699B;A05	ES 185	730266	1827.F15.gz43_232020
M00056699B:G04	ES 185	641072	1827.F21.gz43_232116
M00056699C:C02	ES 185	727649	1827.F24.gz43_232164
M00056699C:F09	ES 185	728196	1827.G02.gz43 231813
M00056699C:G11	ES 185	395604	1827.G04.gz43 231845
M00056699D:C09	ES 186	567122	1827.G07.gz43 231893
M00056700A:F12	ES 186	728300	1827.G14.gz43_232005
M00056700A;G05	ES 186	446607	1827.G16.gz43 232037
M00056700B:D02	ES 186	727151	1827.G17.gz43 232053
M00056700D:H07	ES 186	732770	1827.H10.gz43 231942
M00056701A:A02	ES 186	496084	1827.H12.gz43 231974
M00056701A:G09	ES 186	734264	1827.H16.gz43 232038
M00056701B;D02	ES 186	631056	1827.H18.gz43 232070
M00056701B:D06	ES 186	724489	1827.H19.gz43 232086

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Table 13 CloneID	ES No	ClusterID	SequenceName
M00056701C:F08	ES 186	728196	1827.H24.gz43 232166
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M00056701D:D04	ES 186	727948	1827.I06.gz43_231879
M00056701D:G03	ES 186	553140	1827.107.gz43_231895
M00056701D:G07	ES 186	724059	1827.108.gz43_231911
M00056701D:H12	ES 186	550454	1827,I09.gz43 231927
M00056702C:A10	ES 186	735676	1827.117.gz43 232055
M00056702C:H02	ES 186	448698	1827.123.gz43 232151
M00056702D:G03	ES 186	735834	1827,J04.gz43 231848
M00056703A:G01	ES 186	651050	1827.J09.gz43 231928
M00056703B:A04	ES 186	447624	1827.J13.gz43 231992
M00056703B:D07	ES 186	560678	1827.J16.gz43 232040
M00056703C:F04	ES 186	650136	1827.K01,gz43 231801
M00056703D:F08	ES 186	649927	1827.K06.gz43 231881
M00056704A:B05	ES 186	403671	1827.K08,gz43 231913
M00056704A:H08	ES 186	728552	1827.K13.gz43_231993
M00056704B:C02	ES 186	727760	1827.K14.gz43 232009
M00056704C:C10	ES 186	451183	1827.K18.gz43_232073
M00056704D:H10	ES 186	288801	1827.L01,gz43_231802
M00056705A:B12	ES 186	553805	1827.L03.gz43_231834
M00056705A:D02	ES 186	726852	1827.L04.gz43_231850
M00056705B:A12	ES 186	728768	1827,L09.gz43_231930
M00056705B:D10	ES 186	83388	1827.L10.gz43_231946
M00056706A;B03	ES 186	528981	1827.L24.gz43_232170
M00056706A:B05	ES 186	648777	1827,M01.gz43_231803
M00056706A;E09	ES 186	419465	1827.M02.gz43_231819
M00056706B;C02	ES 186	449752	1827.M06.gz43_231883
M00056706B:C07	ES 186	736100	1827.M07.gz43_231899
M00056706B;D04	ES 186	595181	1827.M08.gz43_231915
M00056706C:A07	ES 186	584693	1827.M10.gz43_231947
M00056706C:B12	ES 186	728898	1827.M11.gz43_231963
M00056706C:D05	ES 186	733064	1827.M13.gz43_231995
M00056706C:G01	ES 186	551450	1827.M18.gz43_232075
M00056707A:G11	ES 186	626791	1827.N03.gz43_231836
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M00056707B:F06	ES 186	736276	1827.N09.gz43_231932
M00056707C;B04	ES 186	427486	1827.N12.gz43_231980
M00056707C:E01	ES 186	552430	1827.N15.gz43_232028
M00056707C:E03	ES 186	730639	1827.N16.gz43_232044
M00056707C:F10	ES 186	728768	1827.N17.gz43_232060
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M00056708B:D03	ES 186	607202	1827.O13.gz43_231997
M00056708B:F06	ES 186	733040	1827.O18.gz43_232077

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M00056708D:H06		M00056708D:B03		728892	
M00056709A:A05	I	M00056708D:D10	ES 186	730463	1827.P06.gz43_231886
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M00056709C:F06		M00056709A:A05	ES 186	454087	
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M00056710B:F05		M00056709C:F06	ES 186		
M00056710B:F05		M00056709D:D05	ES 186	584179	
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M00056710D:A02					
M00056710D:C05					
M00056710-H04		M00056710D:A02			
M00056711A:A09					
M00056711B:B04	[	M00056710D:H04	ES 186	727491	1828.B14.gz43_232384
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M00056713C:A10		M00056713A:F05		48619	
M00056714A:A10		M00056713B:B05	ES 186	727410	
M00056714A:B02         ES 186         735047         1828:E13 gz43 232371           M00056714A:B09         ES 186         728929         1828:E15 gz43 232403           M00056714A:E12         ES 186         448217         1828:E19 gz43 232467           M00056714A:H06         ES 186         728445         1828:E19 gz43 232547           M00056714B:B06         ES 186         728966         1828:E24 gz43 232547           M00056714B:C12         ES 186         724950         1828:F11 gz43 232340           M00056714D:L11         ES 186         729913         1828:F11 gz43 232340           M00056714D:L08         ES 186         50858         1828:F12 gz43 232348           M00056714D:H07         ES 186         50858         1828:F12 gz43 232348           M00056716D:B07         ES 186         558544         1828:G05 gz43 232245           M00056716:B06         ES 186         728182         1828:H02 gz43 232310           M00056716:B06         ES 186         728925         1828:109 gz43 232311           M00056718A:F10         ES 186         728528         1828:109 gz43 232311           M00056718B:C02         ES 186         728528         1828:119 gz43 232347           M00056718C01         ES 186         449585         1828:119 gz43 232347     <					
M00056714A:B09					
M00056714A:E12	1				
M00056714A:H06         ES 186         728445         1828 E22 gz43         232515           M00056714B:B06         ES 186         728966         1828.E24 gz43         232547           M00056714B:C12         ES 186         729913         1828.F11 gz43         232309           M00056714D:A11         ES 186         729913         1828.F11 gz43         232340           M00056714D:B08         ES 186         50858         1828.F12 gz43         232342           M00056714D:H07         ES 186         48217         1828.F14 gz43         232348           M00056715C:B07         ES 186         588544         1828.G05 gz43         232245           M00056716B:F12         ES 186         728182         1828.H02 gz43         23231           M00056716C:B06         ES 186         630516         1828.H04 gz43         23231           M00056718A:F10         ES 186         728925         1828.109 gz43         23231           M00056718A:F0         ES 186         728928         1828.111 gz43         232343           M00056718C:B01         ES 186         451972         1828.H15 gz43         232471           M00056718C:B01         ES 186         646309         1828.H19 gz43         232347           M00056718D:D12	-			1000	
M00056714B:B06					
M00056714B:C12	L				
M00056714D:A11					
M00056714D:E08					
M00056714D:H07         ES 186         448217         1828.F14 gz43 232388           M00056715C:B07         ES 186         558544         1828.G05 gz43 232398           M00056716B:F12         ES 186         558544         1828.G05 gz43 232198           M00056716C:B06         ES 186         728182         1828.H04.gz43 232230           M00056718A:F10         ES 186         728925         1828.H09.gz43 232341           M00056718A:H05         ES 186         728925         1828.111.gz43 232343           M00056718B:C02         ES 186         728924         1828.111.gz43 232447           M00056718C:B01         ES 186         4451972         1828.113.gz43 232471           M00056718D:D12         ES 186         646309         1828.119.gz43 232471           M00056718D:D12         ES 186         449585         1828.123.gz43 232535           M00056719B:D02         ES 186         727946         1828.109.gz43 23212					
M00056715C:B07         ES 186         558544         1828.G05 gz43         232245           M00056716B:F12         ES 186         728182         1828.H02.gz43         232319           M00056716C:B06         ES 186         630516         1828.H04.gz43         232323           M00056718A:F10         ES 186         728925         1828.109 gz43         23231           M00056718A:H05         ES 186         728528         1828.111 gz43         23243           M00056718B:C02         ES 186         451972         1828.H15 gz43         232407           M00056718C:B01         ES 186         646309         1828.H19 gz43         232471           M00056718D:D12         ES 186         449585         1828.L30 gz43         232425           M00056719B:D02         ES 186         1828.H09 gz43         23212 gz43					
M00056716B:F12         ES 186         728182         1828.H02_gz43         232198           M00056716C:B06         ES 186         630516         1828.H04_gz43         232230           M00056718A:F10         ES 186         728925         1828.109_gz43         232311           M00056718A:H05         ES 186         728528         1828.111_gz43         232343           M00056718B:C02         ES 186         451972         1828.115_gz43         232471           M00056718C:B01         ES 186         646309         1828.119_gz43         232471           M00056718D:D12         ES 186         449585         1828.123_gz43         232535           M00056719B:D02         ES 186         727946         1828.109_gz43         23212	١				
M00056716C:B06					
M00056718A:F10         ES 186         72892.5         1828.109 gz43         232311           M00056718A:H05         ES 186         72852.8         1828.11 gz43         232343           M00056718B:C02         ES 186         451972         1828.115 gz43         232407           M00056718C:B01         ES 186         646309         1828.119 gz43         232471           M00056718D:D12         ES 186         449385         1828.123 gz43         232535           M00056719B:D02         ES 186         727946         1828.109 gz43         23212	ı				
M00056718A:H05         ES 186         728528         1828.111 gz43 232343           M00056718B:C02         ES 186         451972         1828.115 gz43 232407           M00056718C:D01         ES 186         646309         1828.119 gz43 232471           M00056718D:D12         ES 186         449585         1828.123 gz43 232535           M00056719B:D02         ES 186         727946         1828.109 gz43 232351					
M00056718B:C02         ES 186         451972         1828:115:gz43         232407           M00056718C:B01         ES 186         646309         1828:119:gz43         2323471           M00056718D:D12         ES 186         449585         1828:123:gz43         232353           M00056719B:D02         ES 186         727946         1828:109:gz43         232312					
M00056718C:B01         ES 186         646309         1828.119.g243_232471           M00056718D:D12         ES 186         449585         1828.123.g243_232535           M00056719B:D02         ES 186         727946         1828.109.g243_232312					
M00056718D:D12 ES 186 449585 1828,123,gz43_232535 M00056719B:D02 ES 186 727946 1828,J09,gz43_232312					
M00056719B:D02 ES 186 727946 1828.J09.gz43_232312					
M00056719B:G04 ES 186 732770 1828.J11.gz43_232344					
	I	M00056719B:G04	ES 186	732770	1828.J11.gz43_232344

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00056719C:B10	ES 186	486363	1828.J18.gz43_232456
M00056719C:F06	ES 186	728303	1828.J19.gz43_232472
M00056720B:D05	ES 186	730233	1828.K06.gz43_232265
M00056721A:C07	ES 186 .	727749	1828.K13.gz43_232377
M00056721B:D03	ES 186	728002	1828.K19.gz43_232473
M00056721C:E05	ES 186	732351	1828.L01.gz43_232186
M00056721C:H01	ES 186	555763	1828.L03.gz43_232218
M00056721D:D01	ES 186	734622	1828.L07.gz43_232282
M00056721D:F12	ES 186	728178	1828,L10.gz43_232330
M00056722A;E10	ES 186	598746	1828.L13.gz43_232378
M00056722A:F08	ES 186	485899	1828.L14.gz43_232394
M00056722B:E09	ES 186	732242	1828.L19.gz43_232474
M00056722C:C09	ES 186	730627	1828.M01.gz43_232187
M00056722C:D11	ES 186	727878	· 1828,M02.gz43_232203
M00056723C:C09	ES 186	462687	1828.N02.gz43_232204
M00056723C:E01	ES 186	553294	1828.N05.gz43_232252
M00056723C:G03	ES 186	734990	1828.N08.gz43 232300
M00056724B:E11	ES 186	473640	1828.N20.gz43 232492
M00056724B:G03	ES 186	447150	1828.N22.gz43 232524
M00056724C:H11	ES 186	726576	1828.O05.gz43_232253
M00056725A:E02	ES 186	728061	1828.O13.gz43 232381
M00056725C:A03	ES 186	727480	1828.O21.gz43_232509
M00056725C:H06	ES 186	728627	1828.P03.gz43_232222
M00056726A:C12	ES 186	735362	1828.P07.gz43_232286
M00056726A:F08	ES 186	449210	1828,P08,gz43_232302
M00056726B:H06	ES 186	447634	1828.P10.gz43_232334
M00056726D:B05	ES 186	692627	1828.P17.gz43_232446
M00056726D:G08	ES 186	509678	1828.P21.gz43_232510
M00056746D:D06	ES 186	729206	1838.B03.gz43_232976
M00056746D:E09	ES 186	458940	1838.B05.gz43_233008
M00056747D:A03	ES 186	529356	1838.C01.gz43_232945
M00056750B:H03	ES 186	735464	1838.E03.gz43_232979
M00056753A:G01	ES 186	552613	1838.G02.gz43_232965
M00056753B:B09	ES 186	728936	1838.G06.gz43_233029
M00056755B:E07	ES 186	728464	1838,I05,gz43_233015
M00056756C:D06	ES 186	736035	1838.J03.gz43_232984
M00056756C:E10	ES 186	646713	1838.J04.gz43_233000
M00056758C:B08	ES 186	730178	1838.K05.gz43_233017
M00056759C:C04	ES 186	736210	1838.L03.gz43_232986
M00056762B:D06	ES 186	481614	1838.N05.gz43_233020
M00056763B:G04	ES 186	697527	1838.O06.gz43_233037
M00056764C:A02	ES 186	551693	1838,P01.gz43_232958
M00056766D:G09	ES 186	728454	1839.A01.gz43_233327
M00056767A:F02	ES 186	730858	1839,A07,gz43_233423

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00056767A:F12	ES 186	711325	1839.A08.gz43_233439
M00056767B:C01	ES 186	727737	1839.A11.gz43_233487
M00056767B:G08	ES 186	736686	1839.A13.gz43_233519
M00056767C:A09	ES 186	549801	1839.A14.gz43_233535
M00056768A:C03	ES 186	730607	1839.A17.gz43_233583
M00056768A;C10	ES 186	728487	1839.A18.gz43_233599
M00056768B:F07	ES 186	730253	1839.A21.gz43_233647
M00056768D:A09	ES 186	736773	1839.B03.gz43_233360
M00056769C:C03	ES 186	504513	1839.B09.gz43_233456
M00056770C:A07	ES 186	641968	1839.B15.gz43_233552
M00056771A:F03	ES 186	732345	1839.B23.gz43_233680
M00056771B:F03	ES 186	552432	1839.C06.gz43_233409
M00056771C:G06	ES 186	640382	1839.C13.gz43_233521
M00056771D:D10	ES 186	726229	1839.C18.gz43_233601
M00056772A:C08	ES 186	691229	1839.C22.gz43_233665
M00056772A:D03	ES 186	726699	1839.C23.gz43_233681
M00056773A:A04	ES 186	735071	1839.D18.gz43_233602
M00056773A:C04	ES 186	735872	1839.D21.gz43_233650
M00056773A:G10	ES 186	727366	1839.D22.gz43_233666
M00056773B:G12	ES 186	711797	1839.E03.gz43_233363
M00056773C:C09	ES 186	510596	1839.E05.gz43_233395
M00056773C:F12	ES 186	135593	1839.E08.gz43_233443
M00056774B:G06	ES 186	726825	1839.E19.gz43_233619
M00056774C:G03	ES 186	728350	1839.E22.gz43_233667
M00056774D:B02	ES 186	631038	1839.E24.gz43_233699
M00056774D:F06	ES 186	734724	1839.F04.gz43_233380
M00056775A:A05	ES 186	727436	1839.F05.gz43_233396
M00056775B:H07	ES 186	477053	1839.F12.gz43_233508
M00056776A:G04	ES 186	728464	1839.G06.gz43_233413
M00056776D:B02	ES 186	523868	1839.G18.gz43_233605
M00056776D:H03	ES 186	642079	1839,G22.gz43_233669
M00056777A;A03	ES 186	730029	1839.G23.gz43_233685
M00056777D:D04	ES 186	730484	1839,H14.gz43_233542
M00056778C:G08	ES 186	609459	1839.H24.gz43_233702
M00056780B:E06	ES 186	724183	1839.J04.gz43_233384
M00056780B;H04	ES 186	493193	1839.J09.gz43_233464
M00056780C:H12	ES 186	647991	1839.J14.gz43_233544
M00056780D:G05	ES 186	727216	1839,J19.gz43_233624
M00056780D:H10	ES 186	724722	1839.J22.gz43_233672
M00056781A:C05	ES 186	513156	1839.J23.gz43_233688
M00056781A:D02	ES 186	482788	1839,J24,gz43_233704
M00056781A:E06	ES 186	730505	1839.K02.gz43_233353
M00056781B:C03	ES 186	733744	1839.K07.gz43_233433
M00056781B:C05	ES 186	639178	1839.K08.gz43_233449

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00056781B:E01	ES 186	<b>728</b> 133	1839.K09.gz43_233465
M00056781B:F05	ES 186	727005	1839.K10.gz43_233481
M00056782B:G05	ES 186	449010	1839,L06.gz43_233418
M00056782B;H11	ES 186	643522	1839.L07.gz43_233434
M00056782D:F02	ES 186	483549	1839.L13.gz43_233530
M00056782D:F08	ES 186	490393	1839.L14.gz43_233546
M00056782D:F10	ES 186	726261	1839.L15.gz43_233562
M00056783A:C08	ES 187	577305	1839.L18.gz43_233610
M00056783B:B01	ES 187	47461	1839.L19.gz43_233626
M00056783C:D01	ES 187	591449	1839.M01.gz43_233339
M00056783C:E03	ES 187	730352	1839,M04.gz43_233387
M00056783D:A02	ES 187	460023	1839.M06.gz43_233419
M00056783D:B07	ES 187	556458	1839,M07.gz43_233435
M00056784B:A01	ES 187	733891	1839.M15.gz43_233563
M00056784B:C02	ES 187	730296	1839.M16.gz43_233579
M00056784B:D06	ES 187	733149	1839.M17.gz43_233595
M00056784C:C06	ES 187	725784	1839.M22.gz43_233675
M00056784D:C06	ES 187	675768	1839.N01.gz43_233340
M00056784D:G10	ES 187	726344	1839.N03.gz43_233372
M00056785A:C12	ES 187	646688	1839.N05.gz43_233404
M00056785A;G06	ES 187	228118	1839.N07.gz43_233436
M00056785B:B07	ES 187	730845	1839.N12.gz43_233516
M00056785C:B09	ES 187	643968	1839.N14.gz43_233548
M00056785D:C09	ES 187	550780	1839.N19.gz43_233628
M00056785D:G07	ES 187	730592	1839.N24.gz43_233708
M00056786D:A03	ES 187	730022	1839.O15.gz43_233565
M00056787B:C07	ES 187	577305	1839.P01.gz43_233342
M00056787C:B04	ES 187	724383	1839.P08.gz43_233454
M00056787C:G01	ES 187	726384	1839.P12.gz43_233518
M00056788A:D06	ES 187	606076	1839.P21.gz43_233662
M00056871C:D05	ES 187	735801	1852.A13.gz43_235742
M00056873A:H06	ES 187	735412	1852.B17.gz43_235807
M00056873B:C09	ES 187	726408	1852.B19.gz43_235839
M00056874C:D05	ES 187	730059	1852,C19.gz43_235840
M00056875D:C04	ES 187	729981	1852.D16.gz43_235793
M00056875D:E09	ES 187	727602	1852.D18.gz43_235825
M00056875D:H12	ES 187	733040	1852.D21.gz43_235873
M00056876A:C08	ES 187	724187	1852.D23.gz43_235905
M00056876A:E02	ES 187	730152	1852,D24.gz43_235921
M00056876C:B02	ES 187	481641	1852.E14.gz43_235762
M00056876C:G10	ES 187	732837	1852.E16.gz43_235794
M00056877B:H09	ES 187	488349	1852.F14.gz43_235763
M00056877C:D11	ES 187	727975	. 1852,F15.gz43_235779

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CloneID	ES No	ClusterID	SequenceName
M00056877C:G12	ES 187	655327	1852.F19.gz43_235843
M00056877C:H03	ES 187	730441	1852.F20.gz43_235859
M00056879A;D12	ES 187	732752	1852.G13.gz43_235748
M00056879B:H11	ES 187	730336	1852.G24.gz43_235924
M00056879D:D01	ES 187	472704	1852.H13.gz43_235749
M00056881A:A10	ES 187	649299	1852,I13.gz43_235750
M00056881A:C02	ES 187	735096	1852.I14.gz43_235766
M00056881A:H02	ES 187	639950	1852.I16.gz43_235798
M00056881B:G04	ES 187	730301	1852.I22.gz43_235894
M00056882B:E12	ES 187	594434	1852.J14.gz43_235767
M00056882D:A06	ES 187	471931	1852.J22.gz43_235895
M00056883D:A07	ES 187	732712	1852.K15.gz43_235784
M00056883D:F07	ES 187	732872	1852.K21.gz43_235880
M00056884C:H08	ES 187	550673	1852.L14.gz43_235769
M00056884C:H11	ES 187	472307	1852.L15.gz43_235785
M00056884D:C07	ES 187	724517	1852.L16.gz43_235801
M00056884D:D06	ES 187	561396	1852.L18.gz43_235833
M00056885A:D12	ES 187	730106	1852,L23,gz43_235913
M00056885D:D01	ES 187	644843	1852.M17,gz43_235818
M00056886A:A09.	ES 187	732872	1852.M19.gz43_235850
M00056886A:C11	ES 187	735326	1852,M21,gz43_235882
M00056886B:B10	ES 187	650944	1852.N13.gz43_235755
M00056886B:C05	ES 187	735672	1852.N14.gz43_235771
M00056886B:H02	ES 187	724223	1852.N17.gz43_235819
M00056886C:D02	ES 187	473512	1852.N19.gz43 235851
M00056886C:D11	ES 187	451184	1852,N20.gz43 235867
M00056887A:E01	ES 187	729087	1852,N23,gz43_235915
M00056887D:H01	ES 187	730195	1852.O17.gz43_235820
M00056888A:H04	ES 187	649030	1852.O24.gz43_235932
M00056890A:D05	ES 187	450963	1861.A06.gz43_236014
M00056891A:G11	ES 187	732315	1861.A24.gz43 236302
M00056891C:H08	ES 187	730393	1861.B08.gz43_236047
M00056892B:C09	ES 187	479572	1861.B18.gz43 236207
M00056893B:G12	ES 187	724773	1861.C16.gz43 236176
M00056893C:A02	ES 187	728640	1861.C17.gz43 236192
M00056895B;F12	ES 187	727596	1861,E03,gz43 235970
M00056895C:E11	ES 187	494393	1861.E07.gz43_236034
M00056895D:G01	ES 187	725258	1861.E14.gz43 236146
M00056896A;C01	ES 187	735834	1861.E18.gz43_236210
M00056898C:B06	ES 187	676221	1861.G10.gz43_236084
M00056898D:D11	ES 187	730760	1861.G17.gz43 236196
M00056898D:H09	ES 187	729060	1861.G20.gz43_236244
M00056899A;A11	ES 187	732676	1861.G22.gz43 236276
M00056899B:D02	ES 187	732244	1861.H04.gz43_235989
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T ADIC 13			
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M00056899D:B06	ES 187	448368	1861.H12.gz43_236117
M00056900B:D01	ES 187	656667	1861.H21.gz43_236261
M00056900B:F07	ES 187	730296	1861.H24.gz43_236309
M00056900C:B12	ES 187	562345	1861.I04.gz43_235990
M00056901B;C03	ES 187	730608	1861.I16.gz43_236182
M00056902A:G12	ES 187	557525	1861.J14.gz43_236151
M00056902D:H09	ES 187	489001	1861.K12.gz43_236120
M00056903A:C08	ES 187	642897	1861.K16.gz43_236184
M00056903A:F03	ES 187	733425	1861.K18.gz43_236216
M00056903B:D09	ES 187	453001	1861.K24.gz43_236312
M00056905A:H04	ES 187	732486	1861.M12.gz43_236122
M00056906D:D11	ES 187	639223	1861.N10.gz43_236091
M00056908B:D02	ES 187	548275	1861.N16.gz43_236187
M00056908D:A11	ES 187	725451	1861.N24.gz43_236315
M00056909B:E11	ES 187	648774	1861.O11.gz43_236108
M00056910A:G05	ES 187	551602	1861.P02.gz43_235965
M00056910B:F01	ES 187	472226	1861.P04.gz43_235997
M00056911B:A01	ES 187	732653	1861.P19.gz43_236237
M00056934A:A12	ES 187	729809	1863,A01,gz43_236702
M00056934A:E07	ES 187	446820	1863.A03.gz43_236734
M00056934C:C04	ES 187	492893	1863.A08.gz43_236814
M00056934C:G09	ES 187	555478	1863.A10.gz43_236846
M00056935A;C02	ES 187	731355	1863.A18.gz43_236974
M00056935A:F04	ES 187	735484	1863,A21,gz43_237022
M00056935C:G08	ES 187	726825	1863.B11.gz43_236863
M00056936C:C06	ES 187	524706	1863.B24.gz43_237071
M00056936C:E04	ES 187	730851	1863.C01.gz43_236704
M00056936C:F11	ES 187	77737	1863.C02.gz43_236720
M00056937C:H08	ES 187	485880	1863.C24.gz43_237072
M00056938A:H09	ES 187	451624	1863.D08.gz43_236817
M00056938B:C06	ES 187	725638	1863.D11.gz43_236865
M00056938B:C09	ES 187	733144	1863.D12.gz43_236881
M00056938B;H08	ES 187	731216	1863.D15.gz43_236929
M00056938C:F10	ES 187	735306	1863.D20.gz43_237009
M00056939A:C01	ES 187	734151	1863.E03.gz43_236738
M00056939B:E05	ES 187	642528	1863.E12.gz43_236882
M00056939B:E10	ES 187	735477	1863.E13.gz43_236898
M00056939D:B02	ES 187	171511	1863.E21.gz43_237026
M00056940B:B11	ES 187	732886	1863.F12.gz43_236883
M00056940B:G07	ES 187	730564	1863 F14.gz43_236915
M00056940C:E05	ES 187	730815	1863.F17.gz43_236963
M00056941B:B02	ES 187	616541	1863.G10.gz43_236852
M00056941B:G08	ES 187	730375	1863.G18.gz43_236980
M00056941C:F07	ES 187	732491	1863.G21.gz43_237028

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00056941D:A05	ES 187	730452	1863.G24.gz43_237076
M00056941D:D07	ES 187	727596	1863.H01.gz43_236709
M00056941D:G05	ES 187	456183	1863.H03.gz43_236741
M00056941D:H01	ES 187	724162	1863.H04.gz43_236757
M00056942B:F04	ES 187	485183	1863.H09.gz43_236837
M00056942C:A12	ES 187	724286	1863.H12.gz43_236885
M00056942C:B12	ES 187	731162	1863.H14.gz43_236917
M00056942C:C06	ES 187	733151	1863.H15.gz43_236933
M00056942C:F11	ES 187	732756	1863.H17.gz43_236965
M00056942D:D03	ES 187	727517	1863.H20.gz43_237013
M00056942D:D07	ES 187	728076	1863.H21.gz43_237029
M00056942D:D11	ES 187	511351	1863.H22.gz43_237045
M00056943A:B04	ES 187	734629	1863.I03.gz43_236742
M00056943A:F01	ES 187	729691	1863.I05.gz43_236774
M00056943C:A01	ES 187	468015	1863.I17.gz43_236966
M00056943D:B02	ES 187	729316	1863.I20.gz43_237014
M00056943D:H08	ES 187	731125	1863.I22.gz43_237046
M00056944A:D09	ES 187	733623	1863.J03.gz43_236743
M00056944B:C02	ES 187	448233	1863.J04.gz43_236759
M00056944C:B03	ES 187	735994	1863.J12.gz43 236887
M00056944D;A06	ES 187	733081	1863.J17.gz43 236967
M00056945A;B11	ES 187	651088	1863.K04.gz43 236760
M00056945A;F02	ES 187	646314	1863.K06.gz43 236792
M00056945C:A11	ES 187	724401	1863.K12.gz43 236888
M00056946A;F07	ES 187	736293	1863.L01.gz43_236713
M00056946A;G06	ES 187	638983	1863.L02.gz43 236729
M00056946C;B08	ES 187	552416	1863.L08.gz43 236825
M00056946D;G09	ES 187	555484	1863.L17.gz43 236969
M00056947A;C05	ES 187	551441	1863.L20.gz43 237017
M00056947C:B04	ES 187	736001	1863,M02.gz43 236730
M00056947D:F09	ES 187	736349	1863,M08,gz43 236826
M00056948A:D09	ES 187	736129	1863,M12.gz43 236890
M00056948B:B03	ES 187	449061	1863.M15.gz43 236938
M00056948B:D04	ES 187	600347	1863.M17.gz43 236970
M00056948B:G05	ES 187	467803	1863,M18.gz43 236986
M00056948B:H06	ES 187	490903	1863.M19.gz43 237002
M00056948C:F03	ES 187	733910	1863.N01.gz43 236715
M00056948D;A11	ES 187	733856	1863,N03,gz43 236747
M00056949A:A04	ES 187	500239	1863,N05,gz43 236779
M00056949D:D11	ES 187	461486	1863.O02.gz43 236732
M00056950A:F01	ES 187	685968	1863.008.gz43 236828
M00056950A:H03	ES 187	410487	1863.O10.gz43 236860
M00056950B:F03	ES 187	560252	1863.O16.gz43 236956
M00056950D:F12	ES 187	731196	1863.P02.gz43 236733
11100030330D.1 12	107	.51150	1000 iz 02.5210_200100

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CloneID	ES No	ClusterID	SequenceName
M00056950D:H09	ES 187	727480	1863.P03.gz43_236749
M00056951B:B06	ES 187	488447	1863.P11.gz43_236877
M00056951B:F09	ES 187	448046	1863.P13.gz43_236909
M00056951C:A04	ES 187	735818	1863.P17.gz43 236973
M00056953B:F05	ES 187	516522	1864.B04.gz43_237135
M00056954C:C04	ES 187	556141	1864.C07.gz43_237184
M00056957B:F10	ES 187	553000	1864.E18.gz43_237362
M00056959C:B10	ES 187	642781	1864.G12.gz43_237268
M00056961C:C07	ES 187	736197	1864.I03.gz43_237126
M00056961D:G02	ES 187	656070	1864.I09.gz43_237222
M00056962D:F09	ES 187	736289	1864.I17.gz43_237350
M00056968C:C06	ES 187	731089	1864.M04.gz43_237146
M00056969A:B07	ES 187	731947	1864.M14.gz43_237306
M00056969A:C07	ES 187	448712	1864.M16.gz43_237338
M00056975A:H11	ES 187	728249	1873.A13.gz43_237662
M00056975B:E04	ES 187	735440	1873.A16.gz43_237710
M00056975C:F09	ES 187	477399	1873.A18.gz43_237742
M00056976C:F05	ES 187	734561	1873.B20.gz43_237775
M00056977A:C02	ES 187	733563	1873.C04.gz43_237520
M00056977A:H10	ES 187	482461	1873.C12.gz43_237648
M00056977B:B06	ES 187	613029	1873.C13.gz43_237664
M00056978A:A03	ES 187	734936	1873.D08.gz43_237585
M00056978A:H04	ES 187	730089	1873.D11.gz43_237633
M00056978B:G02	ES 187	641658	1873.D16.gz43_237713
M00056978D:A01	ES 188	733778	1873.D24.gz43 237841
M00056978D:B07	ES 188	735123	1873.E02.gz43_237490
M00056979B:D03	ES 188	731607	1873.E08.gz43_237586
M00056979B:E03	ES 188	725321	1873.E09.gz43 237602
M00056979C:D11	ES 188	648034	1873.E12.gz43_237650
M00056980A:H06	ES 188	732438	1873.F04.gz43_237523
M00056980D:E07	ES 188	727151	1873.F20.gz43_237779
M00056981D:H02	ES 188	732535	1873.G14.gz43_237684
M00056982D:B12	ES 188	724411	1873.H02.gz43 237493
M00056983D:C12	ES 188	650919	1873.H18.gz43_237749
M00056985A:D06	ES 188	729623	1873.I12.gz43_237654
M00056985A:G07	ES 188	552254	1873.I14.gz43_237686
M00056985B:G03	ES 188	557488	1873.I18.gz43_237750
M00056985C:C06	ES 188	731453	1873.I20.gz43_237782
M00056986A:E09	ES 188	732213	1873.J06.gz43 237559
M00056987A:C02	ES 188	731592	1873.J24.gz43_237847
M00056987C:D08	ES 188	555423	1873.K04.gz43_237528
M00056987D:A09	ES 188	731302	1873.K10.gz43_237624
M00056988A:B09	ES 188	723895	1873.K17.gz43_237736

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CloneID	ES No	ClusterID	SequenceName
M00056988A:F06	ES 188	642649	1873.K20.gz43_237784
M00056988B:A06	ES 188	731358	1873.K22.gz43_237816
M00056988C:D05	ES 188	703978	1873.L04.gz43_237529
M00056988D:F05	ES 188	732113	1873.L10.gz43_237625
M00056989C:H09	ES 188	639934	1873.M03.gz43_237514
M00056989D:A07	ES 188	731476	1873.M05.gz43_237546
M00056989D:F07	ES 188	728013	1873.M10.gz43_237626
M00056989D:H11	ES 188	731317	1873.M12.gz43_237658
M00056990B:B03	ES 188	729795	1873.M18.gz43 237754
M00056990B:H07	ES 188	642288	1873.M20.gz43_237786
M00056990C:E09	ES 188	735395	1873.N02.gz43_237499
M00056990D:C11	ES 188	725825	1873.N06.gz43 237563
M00056991C:H11	ES 188	406931	1873.N18.gz43 237755
M00056992A:E01	ES 188	471883	1873.N22.gz43_237819
M00056993A:B02	ES 188	550973	1873.O12.gz43 237660
M00056993A:B08	ES 188	731542	1873.O13.gz43_237676
M00056993D:C05	ES 188	662617	1873.P02.gz43_237501
M00056993D:F05	ES 188	732114	1873.P05.gz43_237549
M00056994B:H05	ES 188	736401	1873.P18.gz43_237757
M00056994C:B04	ES 188	572426	1873.P20.gz43_237789
M00056994C:C01	ES 188	483676	1873.P22.gz43_237821
M00056995C:H06	ES 188	631251	1874.A15.gz43_238078
M00056995D:C11	ES 188	731686	1874.A19.gz43_238142
M00056996D:A02	ES 188	731317	1874.B08.gz43_237967
M00056997A:H05	ES 188	550016	1874.B18.gz43_238127
M00056997B:C11	ES 188	559053	1874.B20.gz43_238159
M00056997D:B04	ES 188	737083	1874.C06.gz43_237936
M00056998C:B10	ES 188	456236	1874.D01.gz43_237857
M00056998D:H08	ES 188	726408	1874.D08.gz43_237969
M00056999A:G12	ES 188	734708	1874.D12.gz43_238033
M00056999B:D07	ES 188	726696	1874.D16.gz43_238097
M00057000A:A05	ES 188	733932	1874.E03.gz43_237890
M00057001D:F02	ES 188	448202	1874.F16.gz43_238099
M00057003D:F02	ES 188	734629	1874.H16.gz43_238101
M00057004B:D05	ES 188	497233	1874.H21.gz43_238181
M00057006A:G10	ES 188	730662	1874.J17.gz43_238119
M00057007B:G02	ES 188	561993	1874.K16.gz43_238104
M00057007C:A06	ES 188	736413	1874.K17.gz43_238120
M00057009C:B02	ES 188	732792	1874.N01.gz43_237867
M00057011C:H03	ES 188	724810	1874.P07.gz43_237965
M00057011D:F12	ES 188	733836	1874.P13.gz43_238061
M00057012A:D12	ES 188	556637	1874.P17.gz43_238125
M00057012D:G03	ES 188	732441	1875.A03.gz43_238270
M00057012D:G04	ES 188	733571	1875.A04.gz43_238286

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CloneID	ES No	ClusterID	SequenceName
M00057013B:H07	ES 188	481581	1875,A10,gz43 238382
M00057013C:E09	ES 188	726208	1875.A13.gz43 238430
M00057013C:H01	ES 188	733756	1875.A15.gz43 238462
M00057013D:E03	ES 188	558132	1875.A17.gz43 238494
M00057014A:B11	ES 188	512377	1875.A22.gz43 238574
M00057014B;A02	ES 188	556019	1875.B02.gz43 238255
M00057014B:B01	ES 188	478448	1875.B03.gz43 238271
M00057014B:B06	ES 188	451351	1875.B04.gz43 238287
M00057014C:C08	ES 188	728852	1875.B07.gz43_238335
M00057014C:E01	ES 188	733354	1875.B12.gz43_238415
M00057014C:E03	ES 188	730828	1875.B13.gz43_238431
M00057014D:E05	ES 188	619265	1875.B19.gz43_238527
M00057014D:G10	ES 188	733664	1875.B22.gz43_238575
M00057014D;H01	ES 188	557164	1875.B23.gz43_238591
M00057015A:E02	ES 188	732047	1875.C05.gz43_238304
M00057015A:G06	ES 188	487183	1875,C07.gz43_238336
M00057015A:H12	ES 188	732550	1875.C10.gz43_238384
M00057015B:E01	ES 188	484023	1875,C15.gz43_238464
M00057016A;B04	ES 188	734074	1875,C21.gz43_238560
M00057016A:G01	ES 188	494450	1875.C23.gz43_238592
M00057016B:A09	ES 188	733723	1875.D03.gz43_238273
M00057016B:C01	ES 188	554620	1875.D04.gz43_238289
M00057016B:E06	ES 188	734392	1875.D08.gz43_238353
M00057016B;H08	ES 188	474009	1875.D09.gz43_238369
M00057016C:B05	ES 188	733970	1875.D11.gz43_238401
M00057016C:E04	ES 188	731838	1875.D14.gz43_238449
M00057016C:E11	ES 188	689424	1875.D15.gz43_238465
M00057016D:B07	ES 188	703298	1875.D17.gz43_238497
M00057016D;H03	ES 188	640904	1875.D22.gz43_238577
M00057017A:A07	ES 188	452065	1875,D23.gz43_238593
M00057017A:F11	ES 188	570573	1875,E04.gz43_238290
M00057017A:G04	ES 188	538582	1875.E06.gz43_238322
M00057017C:E03	ES 188	559096	1875,E15.gz43_238466
M00057018A:C05	ES 188	734177	1875,E21.gz43_238562
M00057018C:F02	ES 188	732119	1875.F02.gz43_238259
M00057018D:B10	ES 188	734091	1875.F05.gz43_238307
M00057018D:E05	ES 188	557896	1875.F08.gz43_238355
M00057019A:G04	ES 188	626791	1875.F12.gz43_238419
M00057019C;C08	ES 188	455148	1875.F15.gz43_238467
M00057019C:E03	ES 188	558134	1875.F16.gz43_238483
M00057019D:C02	ES 188	649074	1875.F19.gz43_238531
M00057020A:F09	ES 188	639427	1875.F23.gz43_238595
M00057020D:A05	ES 188	483919	
M00057021B;B07	ES 188	477797	1875.G18.gz43_238516

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CloneID	ES No	ClusterID	SequenceName
M00057021D:F10	ES 188	212768	1875.G24.gz43_238612
M00057022B:A04	ES 188	729731	1875.H09.gz43_238373
M00057022B:F03	ES 188	639629	1875.H11.gz43_238405
M00057022B:H04	ES 188	559699	1875.H12.gz43_238421
M00057022C:D12	ES 188	726476	1875.H14.gz43_238453
M00057022D:E10	ES 188	554693	1875.H23.gz43_238597
M00057023A:H09	ES 188	726173	1875.I03.gz43_238278
M00057023D:D05	ES 188	390968	1875.I13.gz43_238438
M00057023D:D08	ES 188	466920	1875.I14.gz43_238454
M00057023D:E11	ES 188	616985	1875.I15.gz43_238470
M00057024A:D08	ES 188	727761	1875.I21.gz43_238566
M00057024B:A03	ES 188	735021	1875.I23.gz43_238598
M00057024B:A11	ES 188	726044	1875.I24.gz43_238614
M00057024B:F07	ES 188	733625	1875.J01.gz43_238247
M00057024C;A01	ES 188	670124	1875.J04.gz43_238295
M00057024C;G08	ES 188	551485	1875.J06.gz43_238327
M00057024D:D12	ES 188	735396	1875.J10.gz43_238391
M00057024D:H11	ES 188	728797	1875.J15.gz43_238471
M00057025C:A08	ES 188	724296	1875.K02.gz43_238264
M00057025C:D11	ES 188	655312	1875.K04.gz43_238296
M00057026C:H11	ES 188	733673	1875.K21.gz43_238568
M00057026D:A05	ES 188	732598	1875.K23.gz43_238600
M00057027B;B11	ES 188	651049	1875.L07.gz43_238345
M00057027B:E04	ES 188	89082	1875.L08.gz43_238361
M00057027B:F06	ES 188	731577	1875.L11.gz43_238409
M00057027D:A12	ES 188	432159	1875.L18.gz43_238521
M00057027D:D07	ES 188	733209	1875.L21.gz43_238569
M00057027D:G03	ES 188	728791	1875.L24.gz43_238617
M00057028B:B11	ES 188	730296	1875.M11.gz43_238410
M00057028D:D09	ES 188	554080	1875.M19.gz43_238538
M00057029A:C08	ES 188	495241	1875.M23.gz43_238602
M00057029B:G10	ES 188	735412	1875.N07.gz43_238347
M00057029D:A06	ES 188	732712	1875.N14.gz43_238459
M00057029D;F01	ES 188	733479	1875.N15.gz43_238475
M00057030B;B03	ES 188	425203	1875.N22.gz43_238587
M00057030B:F01	ES 188	735989	1875.N23.gz43_238603
M00057030C:A05	ES 188	562769	1875.N24.gz43_238619
M00057030C:B03	ES 188	461486	1875.O01.gz43_238252
M00057031A:G09	ES 188	594013	1875.O12.gz43_238428
M00057031A:H02	ES 188	471522	1875.O13.gz43_238444
M00057031B:A01	ES 188	561338	1875.O14.gz43_238460
M00057032A:C01	ES 188	512863	1875.P08.gz43_238365
M00057032A:F12	ES 188	642693	1875.P10.gz43_238397
M00057032D:A04	ES 188	736385	1875.P18.gz43_238525

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CloneID	Table 13			
M00057033C:B03	CloneID	ES No	ClusterID	SequenceName
M00057033D:F10	M00057033A:D08	ES 188	479208	1875.P24.gz43_238621
M00057034B:B01	M00057033C:B03	ES 188	468296	1876.A08.gz43_238734
M00057034C:A11	M00057033D:F10	ES 188	456469	
M00057034C:G12	M00057034B:B01	ES 188	735972	1876.A23.gz43_238974
M00057035C:E10	M00057034C:A11	ES 188	735221	1876.A24.gz43_238990
M00057036A.C07	M00057034C:G12	ES 188	736385	1876.B05.gz43_238687
M00057036E:G08	M00057035C:E10	ES 188	452833	1876.C03.gz43_238656
M00057036D:B01	M00057036A:C07	ES 188	- 557967	1876.C14.gz43_238832
M00057036D:E09	M00057036B:G08	ES 188	483959	1876.C18.gz43_238896
M00057037D-E08	M00057036D:B01	ES 188	729813	1876.C24.gz43_238992
M00057038A-H07	M00057036D:E09	ES 188	554734	1876.D06.gz43_238705
M00057038C:B06	M00057037D:E08	ES 188	727411	1876.D24.gz43_238993
M00057038C.G08	M00057038A:H07	ES 188	473026	1876.E04.gz43_238674
M00057038D-A12	M00057038C:B06	ES 188	728353	1876.E07.gz43_238722
M00057040E:F01	M00057038C:G08	ES 188	454819	1876.E09.gz43_238754
M00057040D:H04	M00057038D:A12	ES 188	735815	1876.E11.gz43_238786
M00057041D:B11	M00057040B:F01	ES 188	483101	1876.F12.gz43_238803
M00057041D:C08	M00057040D:H04	ES 188	736855	1876.F18.gz43_238899
M00057042E:A10	M00057041D:B11	ES 188	546632	1876.G12.gz43_238804
M00057042D:E06   ES 188	M00057041D:C08	ES 188	492627	1876.G13.gz43_238820
M00057042D-G02	M00057042B;A10	ES 188	727321	1876.G22.gz43_238964
M00057043A.G07	M00057042D:E06	ES 188	736881	1876.H06.gz43_238709
M00057043C:H11	M00057042D:G02	ES 188	733772	1876.H08.gz43_238741
M00057044C:B05	M00057043A:G07	ES 188	732315	1876.H12.gz43_238805
M00057045A:C04   ES 188   734466   1876.J05.gz43_238695     M00057045D:D08   ES 188   645803   1876.L20.gz43_238958     M00057046A:F02   ES 188   732159   1876.K03.gz43_238695     M00057046A:G05   ES 188   732159   1876.K03.gz43_238696     M00057046C:E05   ES 188   732042   1876.K14.gz43_238840     M00057047D:E01   ES 188   732042   1876.L11.gz43_238793     M00057047D:E01   ES 188   643933   1876.L20.gz43_238937     M00057047D:E04   ES 188   735054   1876.L23.gz43_238937     M00057048C:E04   ES 188   735054   1876.L23.gz43_238937     M00057048D:E04   ES 188   735054   1876.M12.gz43_238831     M00057049D:F12   ES 188   732562   1876.M10.gz43_238938     M00057049D:F12   ES 188   732246   1876.M02.gz43_238938     M00057049D:F12   ES 188   732246   1876.M07.gz43_2383810     M00057059D:F06   ES 188   732246   1876.M07.gz43_238938     M00057051D:E09   ES 188   466920   1876.O06.gz43_238716     M00057051D:F07   ES 188   732242   1876.C02.gz43_238988     M00057052D:A07   ES 189   708175   1876.P18.gz43_238948     M00057052D:A07   ES 189   708175   1876.P18.gz43_238938     M00057052D:A07   ES 189   708175   1876.P18.gz43_238909     M00057052D:A07   ES 189   708175   1876.P18.gz43_238909     M00057052D:A07   ES 189   708175   1876.P18.gz43_238909     M00057052D:A07   ES 189   708175   1876.P18.gz43_238909	M00057043C:H11	ES 188	638857	1876.H22.gz43_238965
M00057045D-D08   ES 188	M00057044C:B05	ES 188	735871	1876.I16.gz43_238870
M00057046A.F02	M00057045A:C04	ES 188	734466	1876.J05.gz43_238695
M00057046A:G05	M00057045D:D08	ES 188	645803	1876.J20.gz43_238935
M00057046C:E05   ES 188   732042   1876.K14.gz43_238840   M00057047C:C07   ES 188   725024   1876.L11.gz43_238939   M00057047D:E01   ES 188   643933   1876.L20.gz43_238937   M00057047D:H04   ES 188   735054   1876.L23.gz43_238937   M00057048C:E04   ES 188   481293   1876.M10.gz43_238973   M00057048C:H04   ES 188   481293   1876.M10.gz43_2387878   M00057048D:H11   ES 188   735252   1876.M12.gz43_238318   M00057049D:H12   ES 188   732256   1876.M20.gz43_238343   M00057050B:F06   ES 188   732264   1876.N70.gz43_238343   M00057051B:E09   ES 188   732242   1876.005.gz43_238343   M00057051D:F07   ES 188   732242   1876.005.gz43_238343   M00057051D:F07   ES 188   732242   1876.O23.gz43_238988   M00057052D:A07   ES 189   708175   1876.P18.gz43_238909	M00057046A:F02	ES 188	732159	1876.K03.gz43_238664
M00057047C.C07   ES 188   72.5024   1876.L1 gz43 238795     M00057047D.E01   ES 188   643933   1876.L20 gz43 238935     M00057047D.H04   ES 188   735054   1876.L23 gz43 238935     M00057048C.E04   ES 188   481293   1876.M10 gz43 238715     M00057048C.H01   ES 188   481293   1876.M10 gz43 238716     M00057048D.H10   ES 188   732562   1876.M20 gz43 2388718     M00057049D.F12   ES 188   732562   1876.M20 gz43 238938     M00057050B.F06   ES 188   732546   1876.N70 gz43 238341     M00057051B.E09   ES 188   732542   1876.005 gz43 238348     M00057051D.F07   ES 188   732242   1876.003 gz43 238988     M00057052D.A07   ES 189   708175   1876.P18 gz43 238909	M00057046A;G05	ES 188	668731	1876.K05.gz43_238696
M00057047D:E01   ES 188   643933   1876.1.20_gz43_238937     M00057047D:H04   ES 188   735054   1876.1.23_gz43_238985     M00057048C:E04   ES 188   481293   1876.M10_gz43_23878     M00057048C:H11   ES 188   558573   1876.M10_gz43_23878     M00057048D:H10   ES 188   732562   1876.M20_gz43_238938     M00057049D:H12   ES 188   732562   1876.M20_gz43_238938     M0005705D:F06   ES 188   732946   1876.N12_gz43_238938     M00057051B:E09   ES 188   466920   1876.006_gz43_238716     M00057051D:F07   ES 188   732242   1876.006_gz43_238938     M00057052D:A07   ES 189   708175   1876.P18_gz43_238998     M00057052D:A07   ES 189   708175   1876.P18_gz43_238909	M00057046C:E05	ES 188	732042	1876.K14.gz43_238840
M00057047D:H04	M00057047C:C07	ES 188	725024	1876.L11.gz43_238793
M00057048C:E04	M00057047D:E01	ES 188	643933	1876.L20.gz43_238937
M00057048C:H11         ES 188         558573         1876.M12_gz43_238810           M00057048D:H10         ES 188         732562         1876.M20_gz43_238938           M00057049D:F12         ES 188         732246         1876.N07_gz43_238938           M00057050B:F06         ES 188         737087         1876.N14_gz43_238843           M00057051B:E09         ES 188         466920         1876.006_gz43_238716           M00057051D:F07         ES 188         732242         1876.O23_gz43_238988           M00057052D:A07         ES 189         708175         1876.P18_gz43_238909	M00057047D:H04	ES 188	735054	1876.L23.gz43_238985
M00057048DH10         ES 188         732562         1876.M20 gz43_238938           M00057049DF12         ES 188         732246         1876.N70 gz43_238731           M00057050B:F06         ES 188         737087         1876.N14 gz43_238843           M00057051B:E09         ES 188         466920         1876.O26 gz43_238716           M00057051D:F07         ES 188         732242         1876.O23 gz43_238988           M00057052D:A07         ES 189         708175         1876.P18 gz43_238909	M00057048C:E04	ES 188	481293	1876.M10.gz43_238778
M00057048D:H10         ES 188         732562         1876.M20 gz43_238938           M00057049D:F12         ES 188         732246         1876.N7 gz43_238731           M00057050B:F66         ES 188         737087         1876.N14.gz43_238843           M00057051B:E09         ES 188         466920         1876.006_gz43_238716           M00057051D:F07         ES 188         732242         1876.O23_gz43_238988           M00057052D:A07         ES 189         708175         1876.P18_gz43_238909	M00057048C:H11	ES 188	558573	1876.M12.gz43 238810
M00057050B:F06         ES 188         737087         1876.N14.gz43_238843           M00057051B:E09         ES 188         466920         1876.006.gz43_238716           M00057051D:F07         ES 188         732242         1876.023.gz43_238988           M00057052D:A07         ES 189         708175         1876.P18.gz43_238909	M00057048D:H10	ES 188	732562	
M00057051B:E09         ES 188         466920         1876.006 gz43_238716           M00057051D:F07         ES 188         732242         1876.023 gz43_238988           M00057052D:A07         ES 189         708175         1876.P18 gz43_238909	M00057049D:F12	ES 188	732246	1876.N07.gz43 238731
M00057051D:F07 ES 188 732242 1876.O23.gz43_238988 M00057052D:A07 ES 189 708175 1876.P18.gz43_238909	M00057050B:F06	ES 188	737087	1876.N14.gz43 238843
M00057051D:F07 ES 188 732242 1876.O23.gz43_238988 M00057052D:A07 ES 189 708175 1876.P18.gz43_238909	M00057051B:E09	ES 188	466920	1876.Q06.gz43 238716
M00057052D:A07 ES 189 708175 1876.P18.gz43_238909	M00057051D:F07	ES 188	732242	
	M00057052D:A07	ES 189	708175	1876.P18.gz43_238909
	M00057053A:A02	ES 189	645262	1885.A03.gz43_239038

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CloneID	ES No	ClusterID	SequenceName
M00057053A:D11	ES 189	446616	1885.A06.gz43_239086
M00057053C:B10	ES 189	732872	1885.A12.gz43_239182
M00057053C:G04	ES 189	733623	1885.A15.gz43_239230
M00057053D:D01	ES 189	677769	1885.A16.gz43 239246
M00057054A:B12	ES 189	729502	1885.A18.gz43_239278
M00057054A;D01	ES 189	736113	1885.A19.gz43 239294
M00057055B;D07	ES 189	726380	1885.B16.gz43 239247
M00057055B:E10	ES 189	730472	1885.B17.gz43 239263
M00057055B:F05	ES 189	567005	1885.B19.gz43_239295
M00057055C:F01	ES 189	729981	1885.C02.gz43_239024
M00057055C:H07	ES 189	728445	1885.C06.gz43_239088
M00057055D:D11	ES 189	725991	1885.C07.gz43_239104
M00057055D:F05	ES 189	733417	1885.C08.gz43_239120
M00057056B;C06	ES 189	134501	1885.C16.gz43_239248
M00057056B:D05	ES 189	455884	1885.C17.gz43_239264
M00057056B:D11	ES 189	731830	1885.C18.gz43_239280
M00057056B:E09	ES 189	735423	1885.C19.gz43_239296
M00057056B:F01	ES 189	735423	1885.C21.gz43_239328
M00057056C:G03	ES 189	734241	1885.D02.gz43_239025
M00057056D:F11	ES 189	732223	1885.D06.gz43_239089
M00057057A:G01	ES 189	654723	1885.D10.gz43_239153
M00057057B:E06	ES 189	422590	1885.D13.gz43_239201
M00057057B:E07	ES 189	471982	1885.D14.gz43_239217
M00057057B:G03	ES 189	736318	1885.D15.gz43_239233
M00057058D:F09	ES 189	647427	1885.E06.gz43_239090
M00057058D:G08	ES 189	451233	1885.E07.gz43_239106
M00057059A;B04	ES 189	731449	1885,E10,gz43_239154
M00057059B:F03	ES 189	597542	1885.E15.gz43_239234
M00057059C:A04	ES 189	614455	1885.E16.gz43_239250
M00057059C:E10	ES 189	502168	1885.E19.gz43_239298
M00057059C:H02	ES 189	620462	1885.E20.gz43_239314
M00057059D:A09	ES 189	476947	1885.E21.gz43_239330
M00057059D:H09	ES 189	735292	1885,F02.gz43_239027
M00057060A:C10	ES 189	473742	1885,F04,gz43_239059
M00057060B:D07	ES 189	552641	1885.F09.gz43_239139
M00057060B:E06	ES 189	449473	1885.F10.gz43_239155
M00057060C:D05	ES 189	731844	1885.F16.gz43_239251
M00057060D:C09	ES 189	450551	1885.F20.gz43_239315
M00057061A:F09	ES 189	735216	1885.F24.gz43_239379
M00057061A:H10	ES 189	641848	1885.G01 gz43_239012
M00057061B:F01	ES 189	449015	1885.G02.gz43_239028
M00057061B:F05	ES 189	594013	1885.G03.gz43_239044
M00057061B:H02	ES 189	733573	1885.G04.gz43_239060
M00057061C:D04	ES 189	595506	1885.G06.gz43_239092

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00057061D:D03	ES 189	736156	1885.G09.gz43_239140
M00057061D:F05	ES 189	452623	1885.G10,gz43_239156
M00057062B:H04	ES 189	447075	1885.G22.gz43_239348
M00057062D:D04	ES 189	723853	1885.H01.gz43_239013
M00057063A;C08	ES 189	728884	1885.H07.gz43_239109
M00057063B:F06	ES 189	736285	1885.H14.gz43_239221
M00057063C:C04	ES 189	736023	1885.H17.gz43_239269
M00057064B:H10	ES 189	481366	1885.I04.gz43_239062
M00057064C:F11	ES 189	541793	1885.I07.gz43_239110
M00057064D:C09	ES 189	402180	1885.I10.gz43_239158
M00057064D:G09	ES 189	729717	1885.I12.gz43_239190
M00057065B:D12	ES 189	723919	1885,I19.gz43_239302
M00057065C:B07	ES 189	420958	1885.I21.gz43_239334
M00057066A:A09	ES 189	422242	1885.J13.gz43_239207
M00057066B:A04	ES 189	727130	1885.J20.gz43 239319
M00057066C:B02	ES 189	572273	1885.J24.gz43_239383
M00057066D;B03	ES 189	558549	1885.K03.gz43_239048
M00057066D:E01	ES 189	627515	1885,K06,gz43_239096
M00057067B:C11	ES 189	550637	1885.K11.gz43_239176
M00057067B:H04	ES 189	449996	1885.K15.gz43_239240
M00057067C:D04	ES 189	736146	1885.K17.gz43_239272
M00057067C:H09	ES 189	730128	1885.K19.gz43_239304
M00057067D:F03	ES 189	472704	1885,K23,gz43_239368
M00057067D:H06	ES 189	726699	1885.L02.gz43_239033
M00057068A:C10	ES 189	736093	1885.L03.gz43_239049
M00057068A:E07	ES 189	73846	1885.L04.gz43_239065
M00057068A:F05	ES 189	559656	1885.L05.gz43_239081
M00057068A:F07	ES 189	730528	1885.L06.gz43_239097
M00057068A:G05	ES 189	736415	1885.L08.gz43_239129
M00057068D:B03	ES 189	724773	1885.L20.gz43_239321
M00057068D:C09	ES 189	455248	1885.L22.gz43_239353
M00057068D;E05	ES 189	736210	1885.L24.gz43_239385
M00057068D:F04	ES 189	554854	1885.M01.gz43_239018
M00057069A:F09	ES 189	560581	1885.M09.gz43_239146
M00057069A:H08	ES 189	732072	1885.M10.gz43_239162
M00057069B:A08	ES 189	453508	1885.M11.gz43_239178
M00057069B:D07	ES 189	656268	1885.M13.gz43_239210
M00057069B:E07	ES 189	465470	1885.M15.gz43_239242
M00057069D:H09	ES 189	730697	1885.N02.gz43_239035
M00057070A:B07	ES 189	556385	1885.N04.gz43_239067
M00057070C:H10	ES 189	706245	1885.N18.gz43_239291
M00057070D:B08	ES 189	735028	1885.N20.gz43_239323
M00057070D:G03	ES 189	649349	1885.N23.gz43_239371
M00057071A:A10	ES 189	731262	1885.O01.gz43 239020

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Table 13			*
CloneID	ES No	ClusterID	SequenceName
M00057071A:B04	ES 189	624133	1885.O02.gz43_239036
M00057071C:G10	ES 189	469852	1885.O15.gz43_239244
M00057071D:E01	ES 189	447758	1885.O21.gz43_239340
M00057071D:G01	ES 189	732351	1885.O22.gz43_239356
M00057072A.C06	ES 189	456530	1885.P02.gz43_239037
M00057072A:G04	ES 189	448727	1885.P04.gz43_239069
M00057072B:E02	ES 189	422687	1885.P08.gz43_239133
M00057072C;A03	ES 189	452224	1885.P10.gz43_239165
M00057072C;A09	ES 189	735283	1885.P12.gz43_239197
M00057072C:C02	ES 189	734787	1885.P14.gz43_239229
M00057072C:H01	ES 189	631526	1885.P18.gz43_239293
M00057073A:B12	ES 189	620159	1885.P22.gz43_239357
M00057073D:H05	ES 189	732487	1886.A18.gz43_239662
M00057077A:A07	ES 189	630291	1886.C24.gz43_239760
M00057077B:B06	ES 189	639629	1886.D03.gz43_239425
M00057077B:D02	ES 189	726786	1886.D06.gz43_239473
M00057078A:E06	ES 189	450563	1886.D19.gz43_239681
M00057079A:F05	ES 189	556212	1886.E23.gz43_239746
M00057082C:G03	ES 189	368965	1886.H07.gz43_239493
M00057087B:A07	ES 189	733910	1886.L03.gz43_239433
M00057087B:G10	ES 189	727461	1886.L05.gz43_239465
M00057087D:B04	ES 189	734059	1886.L13.gz43_239593
M00057088B:E02	ES 189	731358	1886.M02.gz43_239418
M00057089B:D01	ES 189	538582	1886.M20.gz43_239706
M00057089D:E03	ES 189	731966	1886.M24.gz43_239770
M00057090A:B02	ES 189	473588	1886.N04.gz43_239451
M00057090A:C03	ES 189	727407	1886.N05.gz43_239467
M00057091D:F11	ES 189	732131	1886.O22.gz43_239740
M00057092A:H02	ES 189	729446	1886,P06.gz43_239485
M00057092C:A05	ES 189	419465	1886.P15.gz43_239629
M00057092C:B10	ES 189	729792	1886.P16.gz43_239645
M00057093C:A02	ES 189	553898	1887.A08.gz43_239903
M00057093C:E01	ES 189	736634	1887.A11.gz43_239951
M00057093D:A11	ES 189	557606	1887.A17.gz43_240047
M00057094B:D07	ES 189	533689	1887.B05.gz43_239856
M00057095A:F11	ES 189	735113	1887.B21.gz43_240112
M00057095B:G03	ES 189	732873	1887.C02.gz43_239809
M00057096B:C05	ES 189	477387	1887.C20.gz43_240097
M00057096B:E06	ES 189	730697	1887.C23.gz43_240145
M00057096C:B08	ES 189	731477	1887.D04.gz43_239842
M00057096D:H05	ES 189	457842	1887.D11.gz43_239954
M00057097B:E02	ES 189	729222	1887.D20.gz43_240098
M00057098A:A04	ES 189	560885	1887.E11.gz43_239955
M00057099A:C04	ES 189	644354	1887.F09.gz43_239924

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Table 15			
CloneID	ES No	ClusterID	SequenceName
M00057099A:H12	ES 189	645139	1887.F10.gz43_239940
M00057099B:A06	ES 189	654475	1887.F11.gz43_239956
M00057099B:D02	ES 189	733148	1887.F15.gz43_240020
M00057099B:H11	ES 189	630655	1887.F18.gz43 240068
M00057099C:A06	ES 189	733972	1887.F20.gz43_240100
M00057099D:D11	ES 189	732550	1887.G02.gz43_239813
M00057100C:F02	ES 189	735584	1887.G18.gz43_240069
M00057100C:F03	ES 189	559361	1887.G19.gz43_240085
M00057100D:B06	ES 189	736409	1887.H02.gz43_239814
M00057100D:E09	ES 189	735515	1887.H05.gz43_239862
M00057102A:F12	ES 189	485441	1887.I04.gz43_239847
M00057102B:E12	ES 189	543772	1887.I12.gz43_239975
M00057102D:C11	ES 189	452276	1887.I17.gz43 240055
M00057103A;F11	ES 189	650920	1887.I21.gz43_240119
M00057103A:G01	ES 189	737031	1887.I22.gz43_240135
M00057104A:H06	ES 189	448325	1887.J19.gz43_240088
M00057104D:A02	ES 189	731697	1887.K09.gz43 239929
M00057105C:C06	ES 189	516799	1887.L07.gz43_239898
M00057105C:G02	ES 189	734344	1887.L12.gz43_239978
M00057105D:C10	ES 189	733209	1887.L14.gz43_240010
M00057106A:H04	ES 189	678846	1887.M06.gz43_239883
M00057108B:A12	ES 189	693869	1887.N12.gz43_239980
M00057109A:B02	ES 189	729992	1887.O06.gz43_239885
M00057109A:F11	ES 189	732885	1887.O10.gz43_239949
M00057109A:H09	ES 189	733348	1887.O12.gz43_239981
M00057109C:D10	ES 189	732312	1887.O19.gz43_240093
M00057110A:A03	ES 189	729560	1887.O24.gz43_240173
M00057110A:E12	ES 189	651121	1887.P02.gz43_239822
M00057110C:A04	ES 189	638908	1887.P06.gz43_239886
M00057110C:B09	ES 189	736009	1887.P08.gz43_239918
M00057110D:E12	ES 189	736220	1887.P17.gz43_240062
M00057112A:C12	ES 189	415825	1888.A01.gz43_240175
M00057112D:G08	ES 189	724952	1888.A18.gz43_240447
M00057113A:A08	ES 189	733171	1888.A20.gz43_240479
M00057113B:F03	ES 189	551624	1888.A23.gz43_240527
M00057114C:E07	ES 189	735018	1888.B20.gz43_240480
M00057114C:F08	ES 189	728479	1888.B21.gz43_240496
M00057115B:G06	ES 189	734737	1888.C18.gz43_240449
M00057115C:B11	ES 189	643517	1888.C19.gz43_240465
M00057115D:C10	ES 189	733225	1888.D01.gz43_240178
M00057115D:D06	ES 189	732113	1888.D02.gz43_240194
M00057115D:F06	ES 189	602673	1888,D05,gz43_240242
M00057116A:B06	ES 189	732896	1888.D07.gz43_240274
M00057116C:H09	ES 189	732476	1888.D23.gz43_240530

Table 13

Table 15			
CloneID	ES No	ClusterID	SequenceName
M00057116D:B10	ES 189	474108	1888.D24.gz43_240546
M00057117D:H06	ES 189	733802	1888.E21.gz43_240499
M00057118B:B04	ES 189	602012	1888.F02.gz43_240196
M00057118B:E10	ES 189	553264	1888.F06.gz43_240260
M00057120A:D01	ES 189	555655	1888.G06.gz43_240261
M00057120B:E08	ES 189	730565	1888.G11.gz43_240341
M00057120C:B09	ES 189	453733	1888.G12.gz43 240357
M00057120C:F08	ES 189	609459	1888.G14.gz43 240389
M00057121B:H10	ES 189	553338	1888.H09.gz43_240310
M00057121C:E08	ES 189	552783	1888.H12.gz43_240358
M00057122A:A07	ES 189	639923	1888.H17.gz43_240438
M00057122A:C04	ES 189	647374	1888.H18.gz43 240454
M00057122A:C11	ES 189	733006	1888.H19.gz43_240470
M00057122B:F04	ES 189	656263	1888.H23.gz43_240534
M00057122C:H08	ES 190	418482	1888.I06.gz43_240263
M00057122D:A01	ES 190	733365	1888.I07.gz43_240279
M00057123A:F09	ES 190	732937	1888.I12.gz43_240359
M00057123A:H09	ES 190	729299	1888.I15.gz43_240407
M00057123C:D03	ES 190	569256	1888.I18.gz43_240455
M00057123D:F07	ES 190	734606	1888.I22.gz43_240519
M00057124A:G08	ES 190	736539	1888.J04.gz43_240232
M00057125A:A07	ES 190	726575	1888,J21.gz43_240504
M00057125A;A12	ES 190	449770	1888.J22.gz43_240520
M00057125A:F07	ES 190	727142	1888.J24.gz43_240552
M00057125C:B11	ES 190	736534	1888.K06.gz43_240265
M00057125C:E02	ES 190	730122	1888.K10.gz43_240329
M00057125D:B02	ES 190	542490	1888.K12.gz43_240361
M00057125D:B09	ES 190	729519	1888.K13.gz43_240377
M00057126C:C05	ES 190	662617	1888.K23.gz43_240537
M00057126D:A04	ES 190	568632	1888.L03.gz43_240218
M00057127A:E10	ES 190	553087	1888.L12.gz43_240362
M00057127A:H10	ES 190	735959	1888.L15.gz43_240410
M00057127B:D10	ES 190	454961	1888.L18.gz43_240458
M00057127C:F03	ES 190	447692	1888.L24.gz43_240554
M00057128A:B04	ES 190	556867	1888.M08.gz43_240299
M00057128C:H02	ES 190	735514	1888.M24.gz43_240555
M00057129D:D07	ES 190	440284	1888.N24.gz43_240556
M00057129D:F03	ES 190	451764	1888.O06.gz43_240269
M00057130A:A02	ES 190	736548	1888,O08,gz43_240301
M00057130A:A11	ES 190	288535	1888.O10.gz43_240333
M00057130A:D10	ES 190	736810	1888.O14.gz43_240397
M00057131D:D02	ES 190	726176	1888.P17.gz43_240446
M00057131D:D07	ES 190	735131	1888.P18.gz43_240462

Table 13

CloneID
M00057132C:C04         ES 190         733951         1897.A04_gz43_2406           M00057133A:B07         ES 190         550227         1897.A15_gz43_2407           M00057133C:B02         ES 190         550225         1897.B05_gz43_2406           M00057134B:D06         ES 190         472068         1897.C03_gz43_2405           M00057135A:B02         ES 190         735989         1897.C23_gz43_2409
M00057133A:B07         ES 190         550227         1897.A15 gz43 2407           M00057133C:B02         ES 190         536225         1897.B05 gz43 2405           M00057134B:D06         ES 190         472068         1897.C03 gz43 2405           M00057135A:B02         ES 190         735989         1897.C03 gz43 2405
M00057133C:B02         ES 190         536225         1897.B05.gz43 2406           M00057134B:D06         ES 190         472068         1897.C03.gz43 2405           M00057135A:B02         ES 190         735989         1897.C23.gz43 2409
M00057134B:D06 ES 190 472068 1897.C03.gz43_2405 M00057135A:B02 ES 190 735989 1897.C23.gz43_2409
M00057135A:B02 ES 190 735989 1897.C23.gz43_2409
M00057135A:H03 ES 190 732544 1897.D01 gz43 2405
M00057135D:D01 ES 190 731476 1897.D12.gz43 2407
M00057136A:F01 ES 190 726810 1897.D18.gz43_2408
M00057136B:F06 ES 190 514142 1897.D21.gz43_2408
M00057137A:C01 ES 190 562008 1897.E02.gz43 2405
M00057137A:H12 ES 190 225960 1897.E06.gz43 2406
M00057137B:C07 ES 190 640662 1897.E12.gz43 2407
M00057138A:F02 ES 190 736288 1897.F05.gz43 2406
M00057138B:B02 ES 190 642273 1897.F09.gz43_2406
M00057138B:H02 ES 190 543772 1897.F12.gz43_2407
M00057138C:D06 ES 190 723853 1897.F15.gz43_2407
M00057138D:F03 ES 190 736318 1897.F18.gz43_2408
M00057138D:F10 ES 190 726440 1897.F19.gz43_2408
M00057139A:B10 ES 190 489001 1897.F23.gz43_2409
M00057139A:G08 ES 190 568031 1897.G02.gz43_2405
M00057139A:G12 ES 190 492691 1897.G04.gz43_2406
M00057139D:G07 ES 190 731317 1897.G19.gz43 2408
M00057140B:H01 ES 190 655327 1897.H08.gz43_2406
M00057140D:B03 ES 190 500758 1897.H15.gz43_2407
M00057140D:F02 ES 190 567005 1897.H17.gz43_2408
M00057141A:D09 ES 190 732965 1897.H20.gz43_2408
M00057141A:G06 ES 190 462779 1897.H23.gz43_2409
M00057142A:H07 ES 190 449035 1897.I18.gz43_2408
M00057142B:F06 ES 190 559004 1897.I22.gz43_2409
M00057144B:B07 ES 190 735087 1897.K12.gz43_2407
M00057144B:D04 ES 190 658271 1897.K13.gz43_2407
M00057144B:F10 ES 190 591449 1897.K16.gz43_2408
M00057144B:H10 ES 190 523171 1897.K17.gz43_2408
M00057144C:A02 ES 190 486076 1897.K18.gz43_2408
M00057144C:G04 ES 190 567122 1897.K23.gz43_2409
M00057144C:G08 ES 190 470684 1897.K24.gz43_2409
M00057144D:D03 ES 190 555960 1897.L02.gz43_2405
M00057145B:B07 ES 190 735131 1897.L12.gz43_2407
M00057145B;E06 ES 190 642570 1897.L14.gz43_2407
M00057145C:H03 ES 190 448202 1897.L22.gz43_2409
M00057145D:E01 · ES 190 462779 1897.M02.gz43_2405
M00057146A:H05 ES 190 735801 1897.M12.gz43_2407
M00057146B:C06 ES 190 735283 1897.M13.gz43_2407

Table 13

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CloneID	ES No	ClusterID	SequenceName
M00057146B:H12	ES 190	397515	1897.M17.gz43_240827
M00057146D:F05	ES 190	549461	1897,M24.gz43_240939
M00057147A:B07	ES 190	728413	1897.N03.gz43_240604
M00057147A:E11	ES 190	642936	1897.N06.gz43_240652
M00057147A:H07	ES 190	735729	1897.N08.gz43_240684
M00057147C:B01	ES 190	457846	1897.N12.gz43_240748
M00057147C:G01	ES 190	730190	1897.N14.gz43 240780
M00057147D:H09	ES 190	718314	1897.N22.gz43 240908
M00057148B:G07	ES 190	621226	1897.O05.gz43_240637
M00057148C:B02	ES 190	549285	1897.O09.gz43_240701
M00057148C:C09	ES 190	736318	1897.O12.gz43_240749
M00057149A:A04	ES 190	729843	1897.O18.gz43_240845
M00057149B:B07	ES 190	519378	1897.P03.gz43_240606
M00057149B:B08	ES 190	448212	1897.P04.gz43_240622
M00057149C:H01	ES 190	735756	1897,P11.gz43_240734
M00057150D:C04	ES 190	447003	1898.A03.gz43_242227
M00057150D:F08	ES 190	450724	1898.A05.gz43_242259
M00057151A:B04	ES 190	650297	1898.A06.gz43_242275
M00057152B:H02	ES 190	480307	1898,B05.gz43_242260
M00057152C:C10	ES 190	640603	1898.B10.gz43_242340
M00057152C:C12	ES 190	527789	1898.B12.gz43_242372
M00057153B:A04	ES 190	734484	1898.B22.gz43_242532
M00057153B:D02	ES 190	734808	1898.B24.gz43_242564
M00057153B:G07	ES 190	596809	1898.C02.gz43_242213
M00057153D:H01	ES 190	551654	1898.C15.gz43_242421
M00057154C:F04	ES 190	456816	1898.C21.gz43_242517
M00057154D:H07	ES 190	400314	1898.C23.gz43_242549
M00057155A:E11	ES 190	730661	1898.D04.gz43_242246
M00057155A:G11	ES 190	730341	1898.D07.gz43_242294
M00057155A:H07	ES 190	640563	1898.D08.gz43_242310
M00057155C:B07	ES 190	735993	1898.D12.gz43_242374
M00057155C:G04	ES 190	558477	1898.D15.gz43_242422
M00057155C:H07	ES 190	448510	1898.D16.gz43_242438
M00057155D:E12	ES 190	480142	1898,D22.gz43_242534
M00057156B:D10	ES 190	491933	1898.E06.gz43_242279
M00057156C:E08	ES 190	552972	1898.E10.gz43_242343
M00057156D:F02	ES 190	734928	1898.E17.gz43_242455
M00057157D:H08	ES 190	727608	1898.F05.gz43_242264
M00057158B:C02	ES 190	736030	1898.F10.gz43_242344
M00057158C:C10	ES 190	727976	1898.F17.gz43_242456
M00057158C:G05	ES 190	736790	1898.F19.gz43_242488
M00057159C:E11	ES 190	736228	1898.G11.gz43_242361
M00057160A:C02	ES 190	567122	1898.G16.gz43_242441
M00057160A:F11	ES 190	736940	1898.G18.gz43_242473

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00057160B:G08	ES 190	454073	1898.G22.gz43_242537
M00057160D:A11	ES 190	461316	1898.H04.gz43_242250
M00057161B:D02	ES 190	. 402516	1898.H12.gz43_242378
M00057161B:E07	ES 190	735756	1898.H14.gz43_242410
M00057161B:F04	ES 190	448381	1898.H16.gz43 242442
M00057161D:H08	ES 190	730377	1898.I06.gz43 242283
M00057162B:E10	ES 190	728353	1898.I15.gz43_242427
M00057162C:C01	ES 190	735633	1898.I21.gz43 242523
M00057163A:D11	ES 190	483529	1898.J08.gz43 242316
M00057163A:E09	ES 190	607351	1898.J09.gz43_242332
M00057163C:D06	ES 190	736288	1898.J16.gz43_242444
M00057165C:E01	ES 190	736878	1898.L21.gz43 242526
M00057165C:E08	ES 190	599714	1898.L22.gz43_242542
M00057165C:F05	ES 190	723915	1898.L23.gz43_242558
M00057165D:H05	ES 190	648820	1898.M11.gz43_242367
M00057166B:B07	ES 190	736634	1898.M15.gz43_242431
M00057166B:F05	ES 190	448563	1898.M20.gz43_242511
M00057166B:G06	ES 190	737042	1898.M22.gz43_242543
M00057166C:A10	ES 190	449529	1898.N03.gz43_242240
M00057167A:D08	ES 190	726448	1898.N13.gz43_242400
M00057167C:A05	ES 190	732253	1898.N22.gz43_242544
M00057168B:D02	ES 190	583625	1898.O12.gz43_242385
M00057168C:D03	ES 190	737006	1898.O23.gz43_242561
M00057169A:F05	ES 190	649852	1898.P05.gz43_242274
M00057169A:F06	ES 190	733552	1898.P06.gz43_242290
M00057169C:H10	ES 190	737006	1898.P15.gz43_242434
M00057169C:H12	ES 190	734794	1898.P16.gz43_242450
M00057170A:D06	ES 190	733972	1899.A01.gz43_242579
M00057170A:D11	ES 190	471887	1898.P24.gz43_242578
M00057170A:H04	ES 190	730220	1899.A06.gz43_242659
M00057170B:G01	ES 190	734724	1899.A11.gz43_242739
M00057170C:C01	ES 190	725691	1899.A14.gz43_242787
M00057170D:B08	ES 190	480623	1899.A23.gz43_242931
M00057172A:B02	ES 190	723985	1899.B19.gz43_242868
M00057172A:B04	ES 190	549128	1899.B20.gz43_242884
M00057172A:H06	ES 190	639507	1899.B22.gz43_242916
M00057172D:F08	ES 190	473238	1899.C09.gz43_242709
M00057173A:C07	ES 190	528404	1899.C11.gz43_242741
M00057173C:C07	ES 190	535866	1899.C21.gz43_242901
M00057173D:B12	ES 190	562453	1899.D02.gz43_242598
M00057173D:C07	ES 190	729039	1899.D03.gz43_242614
M00057173D:E04	ES 190	422590	1899.D04.gz43_242630
M00057174B:C06	ES 190	734209	1899.D11.gz43_242742
M00057174B:C11	ES 190	561632	1899.D13.gz43_242774

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CloneID	ES No	ClusterID	SequenceName
M00057174B:F11	ES 190	421794	1899,D15,gz43 242806
M00057174B:G03	ES 190	734744	1899,D16,gz43_242800
M00057174B:G12	ES 190	448770	1899.D17.gz43_242828
M00057174C:G07	ES 190	648221	1899.D21.gz43 242902
M00057175B:G12	ES 190	732625	1899.E08.gz43 242695
M00057175C:D02	ES 190	731370	1899.E13.gz43 242775
M00057175C:F01	ES 190	451812	1899.E16.gz43_242823
M00057175D:A12	ES 190	532904	1899.E17.gz43 242839
M00057175D:E12	ES 190	451383	1899.E22.gz43 242919
M00057176A:A04	ES 190	734692	1899.F01.gz43_242584
M00057176A:C02	ES 190	735235	1899.F03.gz43_242616
M00057176A:H10	ES 190	735764	1899.F07.gz43 242680
M00057176A:H12	ES 190	462986	1899.F08.gz43 242696
M00057176B:B07	ES 190	639779	1899.F11.gz43 242744
M00057176B:F10	ES 190	735605	1899.F16.gz43_242824
M00057176C:H08	ES 190	573733	1899.F19.gz43 242872
M00057176D:A03	ES 190	625988	1899.F20.gz43 242888
M00057177C:D07	ES 190	625988	1899.G12.gz43 242761
M00057177C:G11	ES 190	480142	1899.G13.gz43 242777
M00057177C:H04	ES 190	732429	1899.G15.gz43 242809
M00057177D:G05	ES 190	734726	1899.G22.gz43 242921
M00057179C:B05	ES 190	449548	1899.H05.gz43 242650
M00057179C:G10	ES 190	482512	1899.H08.gz43_242698
M00057179D:B09	ES 190	494423	1899.H09.gz43_242714
M00057180A:H11	ES 190	530883	1899.H15.gz43_242810
M00057180B:C06	ES 190	718314	1899.H18.gz43_242858
M00057180B:D01	ES 190	475872	1899.H19.gz43_242874
M00057180B:F05	ES 190	736280	1899.H21.gz43_242906
M00057180B:G06	ES 190	674526	1899.H23.gz43_242938
M00057180B:H05	ES 190	550588	1899.H24.gz43_242954
M00057180C:F09	ES 190	736309	1899.I06.gz43_242667
M00057180D:C10	ES 191	735789	1899,I09,gz43 242715
M00057180D:G06	ES 191	488447	1899.I11.gz43_242747
M00057181C:D10	ES 191	554221	1899.J01.gz43_242.747
M00057181C:G07	ES 191	560183	1899.J03.gz43_242588
M00057181D:C09	ES 191	735801	1899.J07.gz43_242684
M00057181D:H07	ES 191	452243	1899.J11.gz43_242748
M00057181B:H07	ES 191	737116	1899.J18.gz43_242748
M00057182B:D09	ES 191	736773	1899.J22.gz43 242924
M00057182B:D09	ES 191	642936	1899.K04.gz43_242637
M00057182C:C05	ES 191	373615	1899.K06.gz43_242669
M00057182D:A08	ES 191	477098	1899.K07.gz43 242685
M00057182D:B11	ES 191	736595	1899.K09.gz43_242083
MUUU57182D:B11	JES 191	/36595	1899.K09.gz43_242717

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00057186A B03	ES 191	732244	1899.K14.gz43_242797
M00057186A:E12	ES 191	560568	1899,K17.gz43_242845
M00057186B:E07	ES 191	636748	1899, K22.gz43_242925
M00057189A:A08	ES 191	735871	1899,L05.gz43_242654
M00057189C:G11	ES 191	736354	1899.L20.gz43_242894
M00057189D:B08	ES 191	733623	1899.L22.gz43_242926
M00057189D:G08	ES 191	554683	1899.M02.gz43_242607
M00057191A:H09	ES 191	734031	1899.M07.gz43 242687
M00057191B:C06	ES 191	735514	1899.M09.gz43_242719
M00057191C:D02	ES 191	727523	1899.M14.gz43_242799
M00057191D:B08	ES 191	463402	1899.M17.gz43_242847
M00057191D:G10	ES 191	734928	1899.M22.gz43_242927
M00057192A:B04	ES 191	730805	1899.M23.gz43_242943
M00057192A:D05	ES 191	724061	1899.M24.gz43_242959
M00057192A:D12	ES 191	735426	1899.N01.gz43_242592
M00057192A:F01	ES 191	474346	1899.N02.gz43_242608
M00057192A:H05	ES 191	730106	1899.N04.gz43_242640
M00057192B:D10	ES 191	419711	1899.N07.gz43_242688
M00057192C;B11	ES 191	730899	1899.N10.gz43_242736
M00057192D:C10	ES 191	734724	1899.N13.gz43_242784
M00057192D:G02	ES 191	726786	1899.N16.gz43_242832
M00057192D:G04	ES 191	736402	1899.N17.gz43_242848
M00057193B:C11	ES 191	156329	1899.N21.gz43_242912
M00057193C:F11	ES 191	640635	1899.O06.gz43_242673
M00057193D:F06	ES 191	630516	1899.O09.gz43_242721
M00057194A:B07	ES 191	734061	1899.O11.gz43_242753
M00057194A:C06	ES 191	736049	1899.O13.gz43_242785
M00057194A:F01	ES 191	725089	1899.O16.gz43_242833
M00057194A:H01	ES 191	724061	1899.O18.gz43_242865
M00057194B:E05	ES 191	721768	1899.O21.gz43_242913
M00057194B;H08	ES 191	728061	1899,O24.gz43_242961
M00057194C:A03	ES 191	454812	1899.P01.gz43_242594
M00057194C:F02	ES 191	736861	1899,P05.gz43_242658
M00057194C:F10	ES 191	390968	1899.P07.gz43_242690
M00057194D:C12	ES 191	552055	1899.P10.gz43_242738
M00057196A:E03	ES 191	729173	1899.P15.gz43_242818
M00057196B:H10	ES 191	516729	1899.P17.gz43_242850
M00057197D:H10	ES 191	735797	1900, A14.gz43_243171
M00057198B:C02	ES 191	161489	1900.A20.gz43_243267
M00057199B:B06	ES 191	516729	1900,B08.gz43_243076
M00057199C:H08	ES 191	483061	1900.B12.gz43_243140
M00057200B:D04	ES 191	532904	1900.B22.gz43_243300
M00057200D:E03	ES 191	730308	1900.C04.gz43_243013
M00057201A:H03	ES 191	725905	1900.C12.gz43_243141

Table 13

CloneID	ES No	ClusterID	SequenceName
M00057201B:H10	ES 191	494423	1900.C20.gz43_243269
M00057202A:D05	ES 191	446752	1900.D06.gz43_243046
M00057203B:F08	ES 191	552457	1900.E02.gz43_242983
M00057203C:A09	ES 191	734993	1900.E04.gz43_243015
M00057203D:A09	ES 191	689424	1900.E08.gz43_243079
M00057204A:F11	ES 191	736276	1900.E15.gz43_243191
M00057204C:G06	ES 191	561626	1900.E24.gz43_243335
M00057205D:G06	ES 191	734226	1900.F20.gz43_243272
M00057206A:C06	ES 191	640350	1900.G01.gz43_242969
M00057207A:A07	ES 191	736500	1900.G20.gz43_243273
M00057207B:F06	ES 191	562320	1900.H04.gz43_243018
M00057208B:H08	ES 191	731531	1900.I08.gz43_243083
M00057208C:E12	ES 191	732771	1900.I14.gz43_243179
M00057210A:C12	ES 191	726449	1900.K01.gz43_242973
M00057210C:D09	ES 191	736778	1900.K08.gz43_243085
M00057210C:G07	ES 191	730109	1900.K12.gz43_243149
M00057211A:D01	ES 191	733272	1900.K20.gz43_243277
M00057211B:A08	ES 191	648379	1900.K22.gz43_243309
M00057211B:C09	ES 191	653616	1900.K24.gz43_243341
M00057211B:G01	ES 191	559754	1900.L04.gz43_243022
M00057211C:C06	ES 191	736738	1900.L06.gz43_243054
M00057211C:F05	ES 191	451176	1900.L09.gz43_243102
M00057212B:D05	ES 191	450284	1900.L23.gz43_243326
M00057213C:D06	ES 191	648757	1900.M12.gz43_243151
M00057214A;B05	ES 191	736497	1900.M19.gz43_243263
M00057214B:B12	ES 191	649490	1900.M22.gz43_243311
M00057216C:D12	ES 191	194769	1900.N22.gz43_243312
M00057216D:D05	ES 191	556656	1900.O04.gz43_243025
M00057216D:E12	ES 191	642985	1900.O06.gz43_243057
M00057216D:F10	ES 191	639395	1900.O07.gz43_243073
M00057217A:A12	ES 191	724638	1900.O10.gz43_243121
M00057217B:F12	ES 191	449078	1900.O19.gz43_243265
M00057217D:A06	ES 191	712120	1900.P10.gz43_243122
M00057218C:F10	ES 191	454129	1909.A04.gz43_243395
M00057219A:D05	ES 191	726251	1909.A09.gz43_243475
M00057219A:D07	ES 191	733151	1909.A10.gz43_243491
M00057219A:E11	ES 191	733309	1909.A12.gz43_243523
M00057219A:H11	ES 191	736988	1909.A13.gz43_243539
M00057219B:B10	ES 191	591979	1909.A14.gz43_243555
M00057219B:C06	ES 191	474869	1909.A15.gz43_243571
M00057219D:C02	ES 191	737010	1909.A23.gz43_243699
M00057219D:G11	ES 191	380310	1909.B03.gz43_243380
M00057219D:H04	ES 191	733868	1909.B04.gz43_243396
M00057220A;C06	ES 191	472811	1909.B08.gz43_243460

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Table 13

CloneID	ES No	ClusterID	SequenceName
M00057220A:D09	ES 191	642631	1909.B09.gz43 243476
M00057220B;A06	ES 191	613067	1909.B13.gz43 243540
M00057220B:G01	ES 191	476866	1909.B17.gz43 243604
M00057220B:G10	ES 191	735622	1909.B18.gz43_243620
M00057220C:A08	ES 191	556852	1909.B20.gz43 243652
M00057220C;F08	ES 191	735514	1909.B23.gz43 243700
M00057220D:E06	ES 191	465697	1909.C03.gz43_243381
M00057220D:F02	ES 191	735510	1909.C04.gz43 243397
M00057220D:F06	ES 191	736076	1909.C05.gz43 243413
M00057221A:E10	ES 191	453587	1909,C10,gz43 243493
M00057221B:B01	ES 191	631111	1909.C12.gz43 243525
M00057221B:E11	ES 191	735306	1909.C15.gz43_243573
M00057221C:E07	ES 191	553850	1909.C22.gz43 243685
M00057221C:F02	ES 191	418682	1909.C23.gz43 243701
M00057222B:A06	ES 191	398061	1909.D05.gz43 243414
M00057222D:C10	ES 191	735514	1909.D11.gz43 243510
M00057223A:F06	ES 191	732899	1909.D17.gz43 243606
M00057223B:A07	ES 191	734609	1909.D21.gz43 243670
M00057223B:B04	ES 191	736894	1909.D22.gz43 243686
M00057223B:G01	ES 191	726892	1909.E03.gz43_243383
M00057223C:A01	ES 191	549578	1909.E05.gz43_243415
M00057223C:B01	ES 191	727255	1909.E06.gz43_243431
M00057223D:H03	ES 191	736738	1909.E13.gz43_243543
M00057224A:D07	ES 191	454355	1909.E16.gz43_243591
M00057224B:H02	ES 191	727558	1909.E24.gz43_243719
M00057224C:B02	ES 191	475562	1909.F02.gz43_243368
M00057225A:C08	ES 191	550121	1909.F10.gz43_243496
M00057225A;E03	ES 191	602673	1909.F11.gz43_243512
M00057225C:F09	ES 191	734562	1909.F23.gz43_243704
M00057225C:H07	ES 191	734828	1909.G01.gz43_243353
M00057225D:E01	ES 191	734690	1909.G05.gz43_243417
M00057226A:B04	ES 191	646552	1909.G13.gz43_243545
M00057226A:E09	ES 191	374125	1909.G16.gz43_243593
M00057226C:A09	ES 191	734915	1909.G23.gz43_243705
M00057226C:E05	ES 191	427113	1909.H01.gz43_243354
M00057226C:F05	ES 191	175524	1909.H02.gz43_243370
M00057226C:F12	ES 191	642940	1909.H04.gz43_243402
M00057226C:H10	ES 191	648140	1909.H08.gz43_243466
M00057226D:B03	ES 191	735050	1909.H10.gz43_243498
M00057226D:C10	ES 191	735172	1909.H11.gz43_243514
M00057227B:A05	ES 191	732159	1909.H20.gz43_243658
M00057227B:D12	ES 191	560399	1909.H23.gz43_243706
M00057227B:H04	ES 191	607202	1909.H24.gz43_243722
M00057229A:B03	ES 191	632499	1909.I06.gz43_243435

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Table 13

Table 13			
CloneID	ES No	ClusterID	SequenceName
M00057229B:F11	ES 191	647815	1909.I11.gz43_243515
M00057229B:G11	ES 191	650599	1909.I12.gz43_243531
M00057230B:B07	ES 191	616985	1909.I21.gz43_243675
M00057230B:F10	ES 191	641510	1909.I24.gz43_243723
M00057230C:C05	ES 191	723959	1909.J03.gz43_243388
M00057230C:G12	ES 191	598087	1909.J09.gz43_243484
M00057230D:C05	ES 191	456224	1909.J12.gz43_243532
M00057231A:D04	ES 191	733149	1909.J17.gz43_243612
M00057231A:G04	ES 191	405546	1909.J22.gz43_243692
M00057231B:A01	ES 191	561877	1909.K01.gz43_243357
M00057231C:B04	ES 191	735749	1909.K02.gz43_243373
M00057231C:E06	ES 191	737010	1909.K06.gz43_243437
M00057231C:F12	ES 191	398061	1909.K09.gz43_243485
M00057231C:G04	ES 191	735140	1909.K10.gz43_243501
M00057231D:A04	ES 191	639565	1909.K14.gz43_243565
M00057231D:A05	ES 191	732736	1909.K15.gz43_243581
M00057231D:F10	ES 191	486051	1909.K22.gz43_243693
M00057232B:G02	ES 191	728121	1909.L06.gz43_243438
M00057232D:B03	ES 191	555336	1909.L12.gz43_243534
M00057232D:B05	ES 191	735131	1909.L13.gz43_243550
M00057233A:C04	ES 191	724417	1909.L15.gz43_243582
M00057233B:G04	ES 191	454563	1909.M01.gz43_243359
M00057233D:G12	ES 191	463513	1909.M09.gz43_243487
M00057234B:F06	ES 191	727532	1909.M18.gz43_243631
M00057234C:D11	ES 191	664711	1909.M23.gz43_243711
M00057234D:A12	ES 191	733874	1909.N03.gz43_243392
M00057234D:C09	ES 191	726760	1909.N04.gz43_243408
M00057234D:E04	ES 191	601051	1909.N05.gz43_243424
M00057235B:A07	ES 191	731467	1909.N14.gz43_243568
M00057235C:C08	ES 191	734151	1909.N21.gz43_243680
M00057235C:F03	ES 191	734553	1909.N22.gz43 243696
M00057235D:A05	ES 191	733160	1909.O01.gz43_243361
M00057235D:C03	ES 191	417822	1909.003.gz43_243393
M00057236A:F08	ES 191	450340	1909.O10.gz43 243505
M00057236B;D11	ES 191	725120	1909.O13.gz43 243553
M00057236C:C07	ES 191	453132	1909.O19.gz43_243649
M00057236D:H09	ES 191.	625810	1909.P01.gz43_243362
M00057237B:D10	ES 191	735817	1909.P10.gz43 243506
M00057237D:C11	ES 191	724781	1909.P19.gz43_243650
M00057237D:D09	ES 191	473578	1909.P22.gz43_243698
M00057238A:D07	ES 191	641680	1910.A03.gz43_243763
M00057238B;F05	ES 191	551437	1910.A09.gz43 243859
M00057239B:F05	ES 191	647704	1910.B07.gz43_243828

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00057239C:G07	ES 191	729125	1910.B13.gz43_243924
M00057239D:D01	ES 191	456753	1910.B16.gz43_243972
M00057240A:F03	ES 191	730028	1910.C01.gz43_243733
M00057240B:C01	ES 191	730046	1910.C06.gz43_243813
M00042351A:H03	ES 192	499660	1923.A02.gz43_252631
M00042351C:G01	ES 192	498777	1923.A05.gz43_252679
M00042351C:G11	ES 192	452094	1923.A06.gz43_252695
M00042351D:D02	ES 192	496446	1923.A09.gz43_252743
M00042351D:F08	ES 192	451429	1923.A10.gz43_252759
M00057240C:A06	ES 192	733855	1910.C13.gz43_243925
M00057241B:B04	ES 192	734094	1910.D01.gz43_243734
M00057241D:C04	ES 192	734226	1910.D08.gz43_243846
M00057241D:G01	ES 192	733945	1910.D14.gz43_243942
M00057242A:H11	ES 192	734808	1910.D19.gz43_244022
M00057242B:F07	ES 192	735477	1910.D21.gz43_244054
M00057242C:G12	ES 192	611604	1910.E01.gz43_243735
M00057242D:C07	ES 192	650067	1910.E06.gz43_243815
M00057242D:F10	ES 192	453454	1910.E09.gz43_243863
M00057243A:H03	ES 192	457846	1910.E15.gz43_243959
M00057243C:D01	ES 192	733552	1910.F02.gz43_243752
M00057243C:H11	ES 192	734884	1910.F06.gz43_243816
M00057244C:D08	ES 192	481273	1910.F22.gz43_244072
M00057244C:E06	ES 192	455821	1910.F24.gz43_244104
M00057245A;F03	ES 192	555641	1910.G07.gz43_243833
M00057245B:A08	ES 192	43349	1910.G10.gz43_243881
M00057245B:E02	ES 192	484964	1910.G12.gz43_243913
M00057245D:G02	ES 192	476455	1910.G20.gz43_244041
M00057246A:G11	ES 192	558617	1910.H01.gz43_243738
M00057246C:B12	ES 192	733991	1910.H12.gz43_243914
M00057246D:G09	ES 192	563313	1910.H18.gz43_244010
M00057247C:B11	ES 192	734849	1910.I03.gz43_243771
M00057247C:C11	ES 192	732859	1910.I04.gz43_243787
M00057247C:F10	ES 192	550730	1910.I06.gz43_243819
M00057247D;D10	ES 192	734371	1910.I10.gz43_243883
M00057248A:H10	ES 192	734894	1910.I21.gz43_244059
M00057248B;D11	ES 192	733144	1910.J01.gz43_243740
M00057248C;B08	ES 192	733989	1910.J06.gz43_243820
M00057249A:B05	ES 192	733055	1910.J12.gz43_243916
M00057249A;C06	ES 192	555103	1910.J14.gz43_243948
M00057249C:C07	ES 192	473588	1910.J21.gz43_244060
M00057249D:G03	ES 192	639779	1910.K05.gz43_243805
M00057249D:H09	ES 192	449042	1910.K06.gz43_243821
M00057250B:A03	ES 192	731392	1910.K11.gz43_243901

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00057250B:D04	ES 192	726011	1910.K14.gz43_243949
M00057250C:G02	ES 192	735665	1910.K21.gz43_244061
M00057251A;F02	ES 192	514838	1910.L03.gz43_243774
M00057252A:D10	ES 192	731125	1910.L19.gz43_244030
M00057252B:E04	ES 192	647242	1910.M01.gz43_243743
M00057252C:D08	ES 192	732978	1910.M08.gz43 243855
M00057252D:B10	ES 192	639494	1910.M09.gz43_243871
M00057252D:D04	ES 192	585976	1910.M12.gz43 243919
M00057253A:C05	ES 192	731238	1910.M17.gz43 243999
M00057253A:H10	ES 192	735754	1910.M20.gz43 244047
M00057253B;C06	ES 192	725095	1910.M22.gz43 244079
M00057253C:A06	ES 192	729428	1910.N02.gz43 243760
M00057253C:D11	ES 192	735403	1910.N03.gz43 243776
M00057254A:E12	ES 192	477555	1910.N12.gz43 243920
M00057254A:G07	ES 192	557353	1910.N14.gz43 243952
M00057254B:D04	ES 192	552598	1910.N18.gz43 244016
M00057255C:A07	ES 192	558332	1910.O17.gz43 244001
M00057255C:C10	ES 192	732962	1910.O18.gz43 244017
M00057255D;E02	ES 192	449537	1910.O24.gz43 244113
M00057256B:A05	ES 192	661194	1910.P09.gz43 243874
M00057256D:A11	ES 192	649390	1910.P17.gz43 244002
M00057257A;H10	ES 192	726081	1910.P23.gz43_244098
M00057257B:C11	ES 192	514697	1910.P24.gz43_244114
M00057258C:C09	ES 192	731516	1911.A24.gz43_244483
M00057259D:D11	ES 192	420504	1911.B23.gz43_244468
M00057260A:E05	ES 192	630348	1911.C02.gz43_244133
M00057260A;E11	ES 192	728756	1911.C03.gz43_244149
M00057260C:A04	ES 192	734928	1911.C12.gz43_244293
M00057262B:C03	ES 192	482145	1911.D07.gz43_244214
M00057265C:F03	ES 192	725951	1911.D24.gz43_244486
M00057265D:B12	ES 192	735071	1911.E04.gz43_244167
M00057266C:G12	ES 192	554703	1911.E24.gz43_244487
M00057266D:B12	ES 192	733570	1911.F02.gz43 244136
M00057267A;H04	ES 192	731457	1911,F10.gz43_244264
M00057267C:B12	ES 192	650552	1911.F17.gz43_244376
M00057267C:G09	ES 192	735687	1911.F23.gz43_244472
M00057268A:H05	ES 192	640158	1911.G17.gz43_244377
M00057268B:B03	ES 192	652782	1911.G18.gz43_244393
M00057268D:E04	ES 192	446247	1911.H05.gz43_244186
M00057269C:E01	ES 192	725784	1911.H18.gz43_244394
M00057269C:H06	ES 192	732213	1911.H22.gz43_244458
M00057269D:F02	ES 192	586794	1911.I01.gz43_244123
M00057271A:E04	ES 192	480723	1911.J02.gz43_244140
M00057271B:D09	ES 192	450199	1911.J06.gz43_244204

Table 13

CloneID	ES No	ClusterID	SequenceName
M00057272C;D08	ES 192	728790	1911,J23,gz43 244476
M00057273A:C08	ES 192	630655	1911.K04.gz43 244173
M00057273B:A12	ES 192	735834	1911.K10.gz43 244269
M00057273C:F07	ES 192	736294	1911.K14.gz43 244333
M00057273C:F11	ES 192	482985	1911.K15.gz43 244349
M00057273D:A06	ES 192	729321	1911.K19.gz43 244413
M00057273D:B06	ES 192	474691	1911.K20.gz43 244429
M00057274B:C05	ES 192	735249	1911.L06.gz43 244206
M00057274C:G01	ES 192	735676	1911.L13.gz43 244318
M00057275A:G02	ES 192	456011	1911.L21.gz43_244446
M00057275B:A12	ES 192	546642	1911,L24.gz43_244494
M00057275B:B02	ES 192	648710	1911.M01.gz43_244127
M00057277B:C09	ES 192	736014	1911.M11.gz43_244287
M00057277B:E10	ES 192	642142	1911.M12.gz43_244303
M00057279A:G02	ES 192	736014	1911.N04.gz43_244176
M00057280C:B08	ES 192	734849	1911.O01.gz43_244129
M00057281A:D08	ES 192	450692	1911.O11.gz43_244289
M00057281A:H02	ES 192	456065	1911.O12.gz43_244305
M00057283A:E06	ES 192	736860	1911.P04.gz43_244178
M00057283B:D09	ES 192	736860	1911.P10.gz43_244274
M00057283D:B12	ES 192	734131	1911.P23.gz43_244482
M00057283D:D04	ES 192	552249	1911.P24.gz43_244498
M00057287A:H06	ES 192	730187	1912.B21.gz43_244820
M00057287C:B12	ES 192	734546	1912.C01.gz43_244501
M00057290B:A02	ES 192	728768	1912.D08.gz43_244614
M00057290D:G03	ES 192	558045	1912.D16.gz43_244742
M00057291B:D08	ES 192	732300	1912.D24.gz43_244870
M00057291B:H08	ES 192	730189	1912.E03.gz43_244535
M00057292A:B08	ES 192	639427	1912.E19.gz43_244791
M00057292C:C09	ES 192	73 1910	1912.F03.gz43_244536
M00057293A;H03	ES 192	737109	1912.F12.gz43_244680
M00057293B:H04	ES 192	733856	1912.F16.gz43_244744
M00057299C:A08	ES 192	420402	1912.H10.gz43_244650
M00057299D:E04	ES 192	424672	1912.H15.gz43_244730
M00057300A:A08	ES 192	736449	1912.H17.gz43_244762
M00057300A;B06	ES 192	148201	1912.H18.gz43_244778
M00057301A:A08	ES 192	726430	1912.I11.gz43_244667
M00057302C:D04	ES 192	726874	1912.J18.gz43_244780
M00057305B:B10	ES 192	648738	1912.L05.gz43_244574
M00057309B:E10	ES 192	481930	1912.N06.gz43_244592
M00057314A:A10	ES 192	465528	1921.A01.gz43_244883
M00057314B:H06	ES 192	737114	1921.A07.gz43_244979
M00057314D:E09	ES 192	726692	1921.A16.gz43_245123
M00057314D:F04	ES 192	734894	1921.A17.gz43_245139

Table 13

Table 15			
CloneID	ES No	ClusterID	SequenceName
M00057315D:C06	ES 192	736733	1921.B06.gz43_244964
M00057316A:A08	ES 192	736551	1921.B10.gz43_245028
M00057316A:B04	ES 192	552623	1921.B11.gz43_245044
M00057316A:D09	ES 192	727093	1921.B12.gz43 245060
M00057316C:A06	ES 192	457092	1921.B20.gz43 245188
M00057316D:D02	ES 192	640341	1921.C04.gz43 244933
M00057316D:G04	ES 192	734412	1921.C06.gz43 244965
M00057318A:B02	ES 192	550085	1921.C08.gz43 244997
M00057318B:H05	ES 192	730997	1921.C17.gz43_245141
M00057318C:C12	ES 192	480377	1921.C20.gz43_245189
M00057318C:G04	ES 192	655312	1921.C22.gz43_245221
M00057319D:E06	ES 192	556430	1921.D15.gz43_245110
M00057320A:C03	ES 192	641312	1921.D21.gz43_245206
M00057320A:G08	ES 192	470199	1921.E02.gz43_244903
M00057320B:H03	ES 192	710362	1921.E05.gz43_244951
M00057320D:C02	ES 192	447705	1921.E11.gz43_245047
M00057323A:F01	ES 192	51616	1921.E18.gz43_245159
M00057323B:G04	ES 192	729125	1921.F03.gz43_244920
M00057323B:H08	ES 192	77737	1921.F04.gz43_244936
M00057323D:E10	ES 192	473225	1921.F16.gz43_245128
M00057324A:A09	ES 192	567078	1921.F18.gz43_245160
M00057324A:C08	ES 192	727231	1921.F19.gz43_245176
M00057324B:A06	ES 192	513306	1921.F23.gz43_245240
M00057324B:E04	ES 192	735469	1921.G04.gz43_244937
M00057324C:G05	ES 192	73 1748	1921.G11.gz43_245049
M00057324D:E10	ES 192	. 727013	1921.G13.gz43_245081
M00057324D:H03	ES 192	548635	1921.G14.gz43_245097
M00057325B:B09	ES 192	557693	1921,G17,gz43_245145
M00057325B:D06	ES 192	643005	1921.G18.gz43_245161
M00057325B:H04	ES 192	557451	1921.G24.gz43_245257
M00057325C:F03	ES 192	484974	1921.H03.gz43_244922
M00057326A:E11	ES 192	734522	1921.H10.gz43_245034
M00057326D:B10	ES 192	735054	1921.H18.gz43_245162
M00057328A:F02	ES 192	733723	1921.H21.gz43_245210
M00057328A:F10	ES 192	697006	1921.H22.gz43_245226
M00057328C:F11	ES 192	732969	1921,I06.gz43_244971
M00057328D:G02	ES 192	556490	1921.I10.gz43_245035
M00057331C:E10	ES 192	557177	1921.I21.gz43_245211
M00057331C:F10	ES 192	452936	1921.I22.gz43_245227
M00057331D:D08	ES 192	549552	1921.J01.gz43_244892
M00057332A:C06	ES 192	635849	1921.J06.gz43_244972
M00057332C:F12	ES 192	726692	1921.J14.gz43_245100
M00057333A:D08	ES 192	733464	1921.J19.gz43_245180
M00057333B:F03	ES 192	541793	1921.J24.gz43_245260

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CloneID	ES No	ClusterID	SequenceName
M00057334A:C12	ES 192	726406	1921.K16.gz43_245133
M00057334B:F01	ES 192	734582	1921.K21.gz43_245213
M00057334D:A12	ES 192	735425	1921.L04.gz43 244942
M00057334D:E03	ES 192	732114	1921.L07.gz43 244990
M00057336B;E01	ES 192	727018	1921.L13.gz43 245086
M00057336D;F07	ES 192	448431	1921.L21.gz43_245214
M00057337A:A06	ES 192	727132	1921.L22.gz43_245230
M00057337A:A11	ES 192	482722	1921.L23.gz43_245246
M00057337C:G12	ES 192	554647	1921.M10.gz43_245039
M00057337C:H07	ES 192	496909	1921.M13.gz43 245087
M00057337D:F06	ES 192	614455	1921.M16.gz43_245135
M00057338B:D08	ES 192	643909	1921.M24.gz43_245263
M00057338C:H01	ES 192	466697	1921.N05.gz43_244960
M00057339A:E08	ES 192	551607	1921.N12.gz43 245072
M00057339B;C04	ES 192	477688	1921.N17.gz43 245152
M00057339D:C01	ES 192	448250	1921.004.gz43_244945
M00057339D:H09	ES 192	733146	1921.006.gz43_244977
M00057341D:B09	ES 192	482788	1921.P08.gz43_245010
M00057343D:B10	ES 192	481273	1921.P17.gz43_245154
M00057344A:G07	ES 192	642986	1921.P24.gz43_245266
M00042351D:H05	ES 193	451429	1923.A11.gz43_252775
M00042352A:G05	ES 193	451518	1923.A14.gz43_252823
M00042352A:G09	ES 193	456672	1923.A15.gz43_252839
M00042352B:F03	ES 193	498194	1923.A19.gz43_252903
M00042352B:F10	ES 193	451430	1923.A20.gz43_252919
M00042352C:G01	ES 193	451302	1923.A23.gz43_252967
M00042352D:A11	ES 193	450902	1923.A24.gz43_252983
M00042352D:F11	ES 193	498504	1923.B01.gz43_252616
M00042353B:A11	ES 193	493575	1923.B05.gz43_252680
M00042353B;B02	ES 193	494378	1923.B06.gz43_252696
M00042353D:B08	ES 193	494393	1923.B13.gz43_252808
M00042353D:C06	ES 193	451043	1923.B14.gz43_252824
M00042354B;A07	ES 193	450823	1923.B17.gz43_252872
M00042354C:F04	ES 193	451126	1923.B19.gz43_252904
M00042355A:A12	ES 193	494133	1923.B22.gz43_252952
M00042355A:C03	ES 193	451126	1923.B23.gz43_252968
M00042355A:H09	ES 193	424723	1923,C01.gz43_252617
M00042355B:A05	ES 193	450875	1923, C02.gz43_252633
M00042355B:B07	ES 193	451009	1923.C03.gz43_252649
M00042355B:E10	ES 193	497400	1923.C04.gz43_252665
M00042355C:F02	ES 193	451383	1923.C05.gz43_252681
M00042355C:G09	ES 193 ES 193	451470	1923.C07.gz43_252713

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Table 15			
CloneID	ES No	ClusterID	SequenceName
M00042442A:G04	ES 193	499178	1923.C24.gz43_252985
M00042442B:A07	ES 193	448936	1923.D02.gz43_252634
M00042443B:D03	ES 193	451176	1923.D11.gz43_252778
M00042443D:E01	ES 193	497101	1923.D15.gz43_252842
M00042444A:C10	ES 193	446263	1923.D17.gz43_252874
M00042444A;D04	ES 193	451212	1923.D18.gz43_252890
M00042444A:H06	ES 193	499779	1923.D19.gz43_252906
M00042444C:E02	ES 193	451124	1923.E01.gz43_252619
M00042445A:B04	ES 193	487745	1923.E06.gz43_252699
M00042445C:A08	ES 193	451544	1923.E10.gz43_252763
M00042446B:G02	ES 193	450507	1923.E19.gz43_252907
M00042446D:F04	ES 193	450507	1923.E24.gz43_252987
M00042447C:H10	ES 193	448663	1923.F06.gz43_252700
M00042447D:E04	ES 193	450302	1923.F08.gz43_252732
M00042447D:G10	ES 193	492110	1923.F09.gz43_252748
M00042448B:B02	ES 193	450926	1923.F12.gz43_252796
M00042448B:C04	ES 193	451049	1923.F13.gz43_252812
M00042448C:B07	ES 193	494423	1923.F18.gz43_252892
M00042448C:C09	ES 193	451054	1923.F20.gz43_252924
M00042449A:H10	ES 193	448556	1923.F24.gz43_252988
M00042449C:D10	ES 193	450219	1923.G02.gz43_252637
M00042449D:C06	ES 193	452094	1923.G04.gz43_252669
M00042449D:H11	ES 193	450765	1923.G06.gz43_252701
M00042450A:B01	ES 193	450912	1923.G07.gz43_252717
M00042450A;D02	ES 193	495941	1923.G09.gz43_252749
M00042450C:B09	ES 193	494362	1923.G18.gz43_252893
M00042451D:H10	ES 193	451618	1923.H04.gz43_252670
M00042452C:A09	ES 193	486912	1923.H10.gz43_252766
M00042452C:C10	ES 193	449974	1923.H11.gz43_252782
M00042452D:C04	ES 193	488680	1923.H15.gz43_252846
M00042452D:G06	ES 193	492094	1923.H17.gz43_252878
M00042453A:D12	ES 193	450166	1923.H20.gz43_252926
M00042453C:D12	ES 193	451518	1923.I03.gz43_252655
M00042454A:F02	ES 193	450400	1923.I12.gz43_252799
M00042454D;H10	ES 193	450723	1923.I19.gz43_252911
M00042455A;C06	ES 193	488859	1923.I20.gz43_252927
M00042455A:G12	ES 193	492304	1923.I22.gz43_252959
M00042455B:D05	ES 193	454575	1923.I24.gz43_252991
M00042455B:G09	ES 193	492292	1923.J02.gz43_252640
M00042455C:D11	ES 193	450255	1923.J03.gz43_252656
M00042455C:E04	ES 193	490890	1923.J04.gz43_252672
M00042455D:H08	ES 193	446621	1923,J09.gz43_252752
M00042456A:C08	ES 193	456492	1923.J10.gz43_252768
M00042456A:F08	ES 193	450425	1923.J12.gz43_252800

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00042456C:H07	ES 193	492981	1923.J19.gz43_252912
M00042456D:B06	ES 193	453667	1923.J20.gz43_252928
M00042456D:D07	ES 193	100821	1923.J22.gz43_252960
M00042457D:D02	ES 193	489426	1923.K07.gz43_252721
M00042458B;B08	ES 193	59512	1923.K11.gz43_252785
M00042458B:G05	ES 193	558729	1923.K13.gz43_252817
M00042458C:C07	ES 193	562353	1923.K14.gz43_252833
M00042458C:D08	ES 193	395968	1923.K16.gz43_252865
M00042458C:E06	ES 193	552581	1923.K17.gz43_252881
M00042458D;E06	ES 193	562588	1923.K20.gz43_252929
M00042459A:E04	ES 193	562587	1923.K23.gz43_252977
M00042459B:F03	ES 193	556916	1923.L05.gz43_252690
M00042459B;F12	ES 193	468729	1923.L06.gz43_252706
M00042459C:B11	ES 193	562274	1923.L09.gz43_252754
M00042459C:E06	ES 193	562627	1923.L12.gz43_252802
M00042459C:G02	ES 193	526733	1923.L13.gz43_252818
M00042459D:B07	ES 193	552236	1923.L15.gz43_252850
M00042459D:E03	ES 193	560420	1923.L18.gz43_252898
M00042459D:G04	ES 193	527679	1923.L20.gz43_252930
M00042460B:C06	ES 193	374282	1923.L24.gz43_252994
M00042460B:D10	ES 193	556804	1923.M03.gz43_252659
M00042460B:G12	ES 193	559575	1923.M08.gz43_252739
M00042460C:A02	ES 193	550362	. 1923.M11.gz43_252787
M00042460C:B02	ES 193	452232	1923.M14.gz43_252835
M00042460C:G08	ES 193	562603	1923.M22.gz43_252963
M00042460C:H10	ES 193	562543	1923.M23.gz43_252979
M00042460D:H06	EŞ 193	551755	1923.N05.gz43_252692
M00042516A:A06	ES 193	455439	1923.N08.gz43_252740
M00042516A:A10	ES 193	562115	1923.N09.gz43_252756
M00042516A:C08	ES 193	562060	1923.N10.gz43_252772
M00042516B:E03	ES 193	558559	1923.N13.gz43_252820
M00042516C:C04	ES 193	562399	1923.N18.gz43_252900
M00042516D:B11	ES 193	562275	1923.N22.gz43_252964
M00042516D:C01	ES 193	562398	1923.N23.gz43_252980
M00042516D:H08	ES 193	553312	1923.O01.gz43_252629
M00042516D:H09	ES 193	638444	1923.O02.gz43_252645
M00042517C:B04	ES 193	495074	1923.O05.gz43_252693
M00042517D:A12	ES 193	494300	1923.O09.gz43_252757
M00042517D:G04	ES 193	211273	1923.O11.gz43_252789
M00042518D:A08	ES 193	487522	1923.O18.gz43_252901
M00042519D:F09	ES 193	456024	1923.P02.gz43_252646
M00042519D:H07	ES 193	492982	1923.P04.gz43_252678
M00042520A:F09	ES 193	452833	1923.P10.gz43_252774
M00042520C:E12	ES 193	449104	1923.P16.gz43_252870

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Table 15			
CloneID	ES No	ClusterID	SequenceName
M00043296A:E08	ES 193	497517	1923.P23.gz43_252982
M00043301D:B12	ES 193	450999	1924.C17.gz43_245558
M00043303B:E11	ES 193	450349	1924.D15.gz43_245527
M00043304B:A10	ES 193	450805	1924.E04.gz43_245352
M00043304C:E01	ES 193	496957	1924.E09.gz43_245432
M00043305A:H06	ES 193	204862	1924.E13.gz43_245496
M00043306B:A09	ES 193	493746	1924.E24.gz43_245672
M00043307B:F11	ES 193	498242	1924.F18.gz43_245577
M00043308A:D09	ES 193	496752	1924.G03.gz43_245338
M00043308A:F06	ES 193	451456	1924.G04.gz43_245354
M00043309A:H06	ES 193	499693	1924.G17.gz43_245562
M00043310A:F01	ES 193	498509	1924.H03.gz43_245339
M00043310C:B04	ES 193	495105	1924.H08.gz43_245419
M00043311B:H08	ES 193	499700	1924.H18.gz43_245579
M00043312B:A10	ES 193	494325	1924.H24.gz43_245675
M00043312D:A02	ES 193	494306	1924.I10.gz43_245452
M00043313D:B04	ES 193	492544	1924.J11.gz43_245469
M00043315C:D05	ES 193	495951	1924.K14.gz43_245518
M00043316C:F06	ES 193	498951	1924.L15.gz43_245535
M00043316D:F04	ES 193	479604	1924.L19.gz43_245599
M00043316D:F09	ES 193	553779	1924.L21.gz43_245631
M00043317B:B12	ES 193	494625	1924.M04.gz43_245360
M00043317D:C02	ES 193	451081	1924.M11.gz43_245472
M00043318A:G05	ES 193	454815	1924.M17.gz43_245568
M00043319A:D01	ES 193	562516	1924.N19.gz43_245601
M00043321B:E05	ES 193	490401	1924.P05.gz43_245379
M00043322C:F07	ES 193	446732	1924.P23.gz43_245667
M00043324D:D05	ES 193	451245	1933.A19.gz43_245972
M00043326A:H06	ES 193	500040	1933.B12.gz43_245861
M00043328A:E12	ES 193	496909	1933.C10.gz43_245830
M00043329B:H07	ES 193	492779	1933.C17.gz43_245942
M00043329C:D12	ES 193	453068	1933.C18.gz43_245958
M00043330D:G05	ES 193	412621	1933.D06.gz43 245767
M00043331C:G01	ES 193	491933	1933.D10.gz43 245831
M00043332A:D06	ES 193	454873	1933.D15.gz43 245911
M00043332A:E05	ES 193	490550	1933.D16.gz43 245927
M00043334A:F10	ES 193	491212	1933.E07.gz43 245784
M00043335A;D04	ES 193	450252	1933.E21.gz43 246008
M00043335D;E02	ES 193	450283	1933.F05.gz43_245753
M00043336A:D01	ES 193	496678	1933.F08.gz43 245801
M00043336C:A04	ES 193	450804	1933.F18.gz43_245961
M00043337A:C12	ES 193	495610	1933,G03.gz43 245722
M00043340A:B05	ES 193	494973	1933,H02.gz43_245707
M00043340B:B04	ES 193	453078	1933.H03.gz43_245723

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1 able 15			
CloneID	ES No	ClusterID	SequenceName
M00043340D:C03	ES 193	447057	1933.H09.gz43_245819
M00043342B:H07	ES 193	450727	1933.I02.gz43_245708
M00043342C:G02	ES 193	492629	1933.I06.gz43_245772
M00043342D:D09	ES 193	128697	1933.I09.gz43_245820
M00043343B:B10	ES 193	451307	1933.I15.gz43_245916
M00043343D:H03	ES 193	492887	1933.J04.gz43_245741
M00043344C:C11	ES 193	450027	1933.J15.gz43_245917
M00043346D:E10	ES 193	449299	1933.K07.gz43_245790
M00043348A:D02	ES 193	450129	1933.K16.gz43_245934
M00043349A:C08	ES 193	448927	1933.L01.gz43_245695
M00043351C:A07	ES 193	494271	1933.M05.gz43_245760
M00043353D:E12	ES 193	496870	1933.N10.gz43_245841
M00043354A:C12	ES 193	529733	1933.N13.gz43_245889
M00043354A:E06	ES 193	561712	1933.N14.gz43_245905
M00043354B:C04	ES 193	561480	1933.N18.gz43_245969
M00043354B;F12	ES 193	561834	1933.N20.gz43_246001
M00043354D:C01	ES 193	448110	1933.O05.gz43_245762
M00043354D:G01	ES 193	515328	1933.O10.gz43_245842
M00043354D:G02	ES 193	552445	1933.O11.gz43_245858
M00043354D:H08	ES 193	553594	1933.O12.gz43_245874
M00043355A:D01	ES 193	561593	1933.O18.gz43_245970
M00043355B:B01	ES 193	552687	1933.O23.gz43_246050
M00043355B:D12	ES 193	517346	1933.O24.gz43_246066
M00043355C:G12	ES 193	487106	1933.P08.gz43_245811
M00043356B:E12	ES 193	452586	1934.A03.gz43_253031
M00043356C:F03	ES 193	448230	1934.A06.gz43_253079
M00043356C:H07	ES 193	447429	1934.A07.gz43_253095
M00043357A:G04	ES 193	450583	1934.A11.gz43_253159
M00043357B:B02	ES 193	487893	1934.A12.gz43 253175
M00043357B:D01	ES 193	450189	1934.A13.gz43 253191
M00043358A:D02	ES 193	450242	1934.A21.gz43 253319
M00043358B:D06	ES 193	450122	1934.A23.gz43 253351
M00043358C:F07	ES 193	449061	1934.B04.gz43 253048
M00043359B:A04	ES 193	449795	1934.B09.gz43 253128
M00043359B:D06	ES 193	450193	1934.B11.gz43 253160
M00043359C:C06	ES 193	449994	1934.B14.gz43_253208
M00043360C:E07	ES 193	452719	1934.C01.gz43 253001
M00043360C:F11	ES 194	491177	1934.C03.gz43_253033
M00043360D:D01	ES 194	489368	1934.C05.gz43_253065
M00043361B:F02	ES 194	491402	1934.C08.gz43 253113
M00043361D:A02	ES 194	450080	1934.C10.gz43 253145
M00043362C:E02	ES 194	490846	1934.C18.gz43 253273
M00043363A:G03	ES 194	492022	1934.D02.gz43_253018

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00043363B:A11	ES 194	450756	1934.D04.gz43_253050
M00043363D:F07	ES 194	491507	1934.D12.gz43_253178
M00043364B:H10	ES 194	447355	1934.D21.gz43_253322
M00043364D:H11	ES 194	450756	1934.E05,gz43_253067
M00043365A:C06	ES 194	449956	1934.E08.gz43_253115
M00043365B:A10	ES 194	486833	1934.E11.gz43_253163
M00043365B:C08	ES 194	488613	1934.E13.gz43_253195
M00043365C:A09	ES 194	449780	1934.E16.gz43_253243
M00043365C:F06	ES 194	491240	1934.E19.gz43_253291
M00043366A;B12	ES 194	488023	1934.E24.gz43_253371
M00043366D:B07	ES 194	488310	1934.F09.gz43_253132
M00043367C:D10	ES 194	450147	1934.F14.gz43_253212
M00043368A:F03	ES 194	491492	1934.F16.gz43_253244
M00043369D:B01	ES 194	449842	1934.G02.gz43_253021
M00043369D:G10	ES 194	450551	1934,G07.gz43 253101
M00043370A;C08	ES 194	449959	1934.G08.gz43_253117
M00043370B:D08	ES 194	450207	1934.G14.gz43_253213
M00043371A:D06	ES 194	490060	1934.G19.gz43_253293
M00043371B:C10	ES 194	450080	1934,G22.gz43_253341
M00043371C:B02	ES 194	449908	1934.G23.gz43_253357
M00043372B:A07	ES 194	43013	1934.H05.gz43_253070
M00043372B:B01	ES 194	455707	1934,H06.gz43_253086
M00043373A:G11	ES 194	452710	1934,H19,gz43_253294
M00043373B:G12	ES 194	450658	1934.H24.gz43_253374
M00043373D:G04	ES 194	456492	1934.I04.gz43_253055
M00043374A:E05	ES 194	560213	1934.I08.gz43_253119
M00043374A:G04	ES 194	510272	1934.I10.gz43_253151
M00043374D:D07	ES 194	631526	1934.I15.gz43_253231
M00043374D:H09	ES 194	515423	1934.I17.gz43_253263
M00043375A:E01	ES 194	451292	1934.I20.gz43_253311
M00043375A:E02	ES 194	451294	1934.I21.gz43_253327
M00043375B:A04	ES 194	456103	1934.I24.gz43_253375
M00043376B:C03	ES 194	451144	1934.J07.gz43_253104
M00043376B:F06	ES 194	450482	1934.J08.gz43_253120
M00043376D:A04	ES 194	493622	1934.J11.gz43 253168
M00043376D;A12	ES 194	233814	1934.J12.gz43 253184
M00043376D:D12	ES 194	451185	1934.J14.gz43 253216
M00043377C:A11	ES 194	· 449959	1934.J20.gz43_253312
M00043377D:E01	ES 194	451361	1934.J22.gz43_253344
M00043378B:B05	ES 194	494450	1934.J24.gz43_253376
M00043378D:D12	ES 194	496084	1934.K05.gz43_253073
M00043379C:B09	ES 194	451011	1934.K13.gz43_253201
M00043379D:A05	ES 194	494099	1934,K16,gz43_253249
M00043380D:E10	ES 194	450193	1934.L05.gz43_253074

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Table 13	I roar	C1 . TT	
CloneID	ES No	ClusterID	SequenceName
M00043380D:E11	ES 194	451326	1934.L06.gz43_253090
M00043381B;E10	ES 194	551380	1934.L14.gz43_253218
M00043381C:D08	ES 194	561641	1934.L16.gz43_253250
M00043381C:E10	ES 194	556511	1934.L18.gz43_253282
M00043381D:F09	ES 194	467035	1934.L24.gz43_253378
M00043382C:D07	ES 194	496586	1934.M03.gz43_253043
M00043382C:G09	ES 194	499240	1934.M05.gz43_253075
M00043383B:F12	ES 194	456367	1934.M11.gz43_253171
M00043383D;C07	ES 194	495143	1934.M14.gz43_253219
M00043383D:G07	ES 194	451467	1934.M15.gz43_253235
M00043384B:A04	ES 194	487437	1934.M18.gz43_253283
M00043384C:A07	ES 194	487448	1934.M20.gz43_253315
M00043384C:C02	ES 194	489207	1934.M22.gz43_253347
M00043384C:G01	ES 194	492627	1934.M24.gz43_253379
M00043384D:D05	ES 194	489506	1934.N02.gz43_253028
M00043385A:E01	ES 194	496760	1934.N06.gz43_253092
M00043385A:G12	ES 194	448448	1934.N08.gz43_253124
M00043385C;D06	ES 194	496234	1934.N11.gz43_253172
M00043386A:A11	ES 194	450819	1934.N16.gz43_253252
M00043387A:E02	ES 194	490393	1934.O03.gz43_253045
M00043387A:E03	ES 194	490395	1934.O04.gz43_253061
M00043387B:A03	ES 194	486856	1934.O09.gz43 253141
M00043387C:A11	ES 194	487183	1934.O14.gz43_253221
M00043387C:G08	ES 194	455460	1934.O18.gz43_253285
M00043387D:A02	ES 194	449802	1934.O21.gz43_253333
M00043387D:B02	ES 194	488030	1934.O22.gz43_253349
M00043387D:D06	ES 194	450211	1934.O23.gz43_253365
M00043388A:D05	ES 194	453893	1934.P02.gz43_253030
M00043388B:C02	ES 194	489275	1934.P04.gz43_253062
M00043389D:B10	ES 194	488349	1934.P16.gz43_253254
M00043390B:C04	ES 194	449978	1934.P20.gz43_253318
M00043391D:D05	ES 194	217042	1935.A19.gz43_246512
M00043392C:F02	ES 194	491448	1935.B05.gz43 246289
M00043397B:H07	ES 194	335078	1935.D06.gz43 246307
M00043397C:G02	ES 194	450623	1935.D08.gz43_246339
M00043401A:E09	ES 194	490805	1935.E18.gz43 246500
M00043402C:F12	ES 194	456213	1935.F12.gz43 246405
M00043403B:A12	ES 194	450335	1935.F18.gz43 246501
M00043406D:C04	ES 194	448924	1935.H08.gz43 246343
M00043407C:H08	ES 194	451456	1935.H15.gz43_246455
M00043407D:G06	ES 194	498629	1935.H17.gz43 246487
M00043409C;C07	ES 194	402070	1935.I07.gz43 246328
M00043410C:C05	ES 194	451781	1935,I17.gz43 246488
M00043501A:D07	ES 194	553890	1935.K06.gz43 246314

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00043501D:C01	ES 194	562378	1935.K19.gz43_246522
M00043502A:H01	ES 194	491644	1935.L06.gz43_246315
M00043503C:F08	ES 194	402823	1935.M10.gz43_246380
M00043504A:E06	ES 194	477674	1935.M23.gz43_246588
M00043504B:C02	ES 194	635439	1935.N06.gz43_246317
M00043504C:E03	ES 194	555399	1935.N16.gz43 246477
M00043504D:G08	ES 194	446964	1935.N22.gz43 246573
M00043505B:G03	ES 194	562886	1935.O08.gz43_246350
M00043505B:G07	ES 194	558055	1935.O09.gz43_246366
M00043506D:F06	ES 194	522703	1935.P24.gz43_246607
M00043508A:H02	ES 194	456742	1936.B10.gz43_246753
M00043508B:G11	ES 194	562823	1936.B18.gz43_246881
M00043508D:C01	ES 194	414739	1936.C08.gz43_246722
M00054486B:E09	ES 194	562382	1936.D18.gz43_246883
M00054486B:H01	ES 194	549757	1936.D21.gz43_246931
M00054487C:A01	ES 194	559574	1936.E21.gz43_246932
M00054488A:F01	ES 194	556768	1936.F13.gz43_246805
M00054488C:D04	ES 194	564967	1936.G03.gz43_246646
M00054492A:D04	ES 194	449978	1936.I15.gz43_246840
M00054493C:E04	ES 194	489368	1936.J16.gz43_246857
M00054494A:H04	ES 194	549853	1936.K05.gz43_246682
M00054494D:A10	ES 194	560188	1936.K13.gz43_246810
M00054497B:C10	ES 194	560275	1936.M05.gz43_246684
M00054497C:G12	ES 194	450037	1936.M10.gz43_246764
M00054497D:A04	ES 194	554908	1936.M11.gz43_246780
M00054498C:B08	ES 194	558500	1936.N04.gz43_246669
M00054498D:F01	ES 194	449000	1936.N11.gz43_246781
M00054500A:F04	ES 194	559720	1936.O08.gz43_246734
M00054501A:E11	ES 194	552920	1936.O23.gz43_246974
M00054501A:G12	ES 194	488171	1936.O24.gz43_246990
M00054501C:E08	ES 194	481057	1936.P05.gz43_246687
M00054502 C:E02	ES 194	550571	1936.P24.gz43_246991
M00054502C:H10	ES 194	449776	1945.A02.gz43_248938
M00054503D:E07	ES 194	562019	1945.B03.gz43_248955
M00054504B:A04	ES 194	524721	1945.B11.gz43_249083
M00054504D:E02	ES 194	551475	1945.B21.gz43_249243
M00054504D:G07	ES 194	289201	1945.B23.gz43_249275
M00054505A:G12	ES 194	572807	1945.C05.gz43_248988
M00054506A:D05	ES 194	554839	1945.C24.gz43_249292
M00054506B:A07	ES 194	388055	1945.D04.gz43_248973
M00054506B:H01	ES 194	550986	1945.D11.gz43_249085
M00054507B:A07	ES 194	505907	1945.E05.gz43_248990
M00054508B:E08	ES 194	557681	1945.F12.gz43_249103
M00054508C:B04	ES 194	550129	1945.F15.gz43_249151

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00054508C:C08	ES 194	555771	1945.F17.gz43_249183
M00054509A;C01	ES 194	547866	1945.F23.gz43_249279
M00054509B:F01	ES 194	559127	1945.G08.gz43_249040
M00054509C:G01	ES 194	499696	1945.G11.gz43_249088
M00054510C:G07	ES 194	452471	1945.H07.gz43 249025
M00054511A:D11	ES 194	570573	1945.H15.gz43_249153
M00054511B:C06	ES 194	389065	1945.H19.gz43_249217
M00054512D:H08	ES 194	487658	1945,I24,gz43_249298
M00054513A:B08	ES 194	550063	1945.J03.gz43_248963
M00054513A:F09	ES 194	455814	1945.J05.gz43_248995
M00054513B:E11	ES 194	551518	1945.J07.gz43_249027
M00054513C:A01	ES 194	555660	1945.J09.gz43_249059
M00054513D:F04	ES 194	550678	1945.J13.gz43_249123
M00054514C:B01	ES 194	461313	1945.J21.gz43_249251
M00054515B:E11	ES 194	554246	1945.K13.gz43_249124
M00054516C:A04	ES 194	559343	1945.L14.gz43_249141
M00054516D:F09	ES 194	554456	1945.L21.gz43_249253
M00054517A:C01	ES 194	488023	1945.L23.gz43_249285
M00054517B:B04	ES 194	549129	1945.M05.gz43_248998
M00054518C:F11	ES 194	549581	1945.N05.gz43_248999
M00054518D:D03	ES 194	552905	1945.N09.gz43_249063
M00054519A:C04	ES 194	554611	1945.N11.gz43_249095
M00054521D:H01	ES 194	572992	1945.O18.gz43_249208
M00054523C;A11	ES 194	458974	1945.P23.gz43_249289
M00054523D:D07	ES 194	558497	1946.A06.gz43_249386
M00054523D:E08	ES 194	550022	1946.A08.gz43_249418
M00054523D:G09	ES 194	454336	1946,A10,gz43_249450
M00054525D:H05	ES 194	559676	1946.B07.gz43 249403
M00054526A:E04	ES 194	550475	1946,B11.gz43_249467
M00054526B:C09	ES 194	556809	1946.B15.gz43_249531
M00054526C:F03	ES 194	524966	1946.B19.gz43_249595
M00054527B:A07	ES 194	552006	1946.C06.gz43 249388
M00054529A;H01	ES 194	585380	1946.D21.gz43_249629
M00054529C:D11	ES 194	551157	1946.E06.gz43 249390
M00054529D:E01	ES 194	490308	1946,E10,gz43 249454
M00054530D:C10	ES 194	312036	1946.E23.gz43 249662
M00054530D:H07	ES 194	553002	1946.E24.gz43 249678
M00054531D:F05	ES 194	584071	1946,F11.gz43_249471
M00054532A:A07	ES 194	549919	1946.F13.gz43_249503
M00054532B:C05	ES 194	550204	1946.F21.gz43_249631
M00054532D:A01	ES 194	559574	1946.G06.gz43_249392
M00054532D:E07	ES 194	526984	1946.G08.gz43_249424
M00054533A:F05	ES 194	571899	1946.G12.gz43_249488
M00054533B:F04	ES 194	550714	1946.G20.gz43 249616

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00054533C:B04	ES 194	556940	1946.H01.gz43_249313
M00054533C:E07	ES 194	466020	1946.H05.gz43_249377
M00054534A:B06	ES 194	498433	1946.H15.gz43_249537
M00054534D:D02	ES 194	556216	1946.I06.gz43_249394
M00054535A:G03	ES 194	553338	1946.I14.gz43_249522
M00054535B:H08	ES 194	560205	1946.I22.gz43_249650
M00054536B:A03	ES 194	550018	1946.J07.gz43_249411
M00054536B:D02	ES 194	550370	1946.J10.gz43_249459
M00054536B:F08	ES 194	550730	1946.J11.gz43_249475
M00054536C:B11	ES 194	555019	1946.J15.gz43_249539
M00054536C:D12	ES 195	82864	1946.J17.gz43_249571
M00054537B:H03	ES 195	451778	1946.K02.gz43_249332
M00054537C:B03	ES 195	51103	1946.K06.gz43_249396
M00054538A:E10	ES 195	455821	1946.K16.gz43_249556
M00054538B:B05	ES 195	550172	1946.K18.gz43_249588
M00054538D:F05	ES 195	557339	1946.L07.gz43_249413
M00054538D:G12	ES 195	550830	1946.L11.gz43_249477
M00054538D:H11	ES 195	560014	1946.L13.gz43_249509
M00054539B:D06	ES 195	560519	1946.L17.gz43_249573
M00054539B:G03	ES 195	549734	1946.L19.gz43_249605
M00054540A:H07	ES 195	464154	1946.M10.gz43_249462
M00054541B:A09	ES 195	549233	1946.M22.gz43_249654
M00054541C:C10	ES 195	493575	1946.N03.gz43_249351
M00054541C:F11	ES 195	481594	1946.N09.gz43_249447
M00054542B:B01	ES 195	409262	1946.N17.gz43_249575
M00054542C:B12	ES 195	558024	1946.N21.gz43_249639
M00054543B:E06	ES 195	552638	1946.O12.gz43_249496
M00054543C:F01	ES 195	552753	1946.O18.gz43_249592
M00054543D:G08	ES 195	549131	1946.P01.gz43_249321
M00054544B;E03	ES 195	550618	1946.P09.gz43_249449
M00054544C:F04	ES 195	560717	1946.P17.gz43_249577
M00054545C:B09	ES 195	555502	1947.A10.gz43_253539
M00054545D:E04	ES 195	561406	1947.A14.gz43_253603
M00054553D:E09	ES 195	515707	1947.G22.gz43_253737
M00054555A:H09	ES 195	550952	1947.I06.gz43_253483
M00054555D:C03	ES 195	550246	1947.J01.gz43_253404
M00054566D:G08	ES 195	496586	1947.P21.gz43_253730
M00054567C:B03	ES 195	556065	1948.A04.gz43_249738
M00054567C:D09	ES 195	550403	1948.A07.gz43_249786
M00054568A:G02	ES 195	549716	1948.A14.gz43_249898
M00054568A:H03	ES 195	549858	1948.A16.gz43_249930
M00054568B:A07	ES 195	554793	1948.A17.gz43_249946
M00054568C:D06	ES 195	450963	1948.A21.gz43_250010
M00054568C:D06	ES 195	450963	

Table 13

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M00054582A:A05 ES 195 M00054582A:A07 ES 195 M00054583A:B04 ES 195 M00054583A:F05 ES 195 M00054583D:E04 ES 195	556336	1948.M12.gz43_249878
M00054582A:A07 ES 195 M00054583A:B04 ES 195 M00054583A:F05 ES 195 M00054583D:E04 ES 195	550811	1948.M13.gz43_249894
M00054583A:B04 ES 195 M00054583A:F05 ES 195 M00054583D:E04 ES 195	568467	1948.M14.gz43_249910
M00054583A:F05 ES 195 M00054583D:E04 ES 195	549994	1948.M15.gz43_249926
M00054583D:E04 ES 195	550106	1948.N13.gz43_249895
	550694	1948.N14.gz43 249911
	394567	1948.O01.gz43 249704
M00054584A:A07 ES 195	549956	1948.O03.gz43 249736
M00054584A:B03 ES 195	533812	1948.O05.gz43 249768
M00054584B:A03 ES 195	554887	1948.O10.gz43 249848
M00054584B:G03 ES 195	572249	1948.O13.gz43_249896
M00054584D:C01 ES 195		1948.O17.gz43_249960
M00054585A:E07 ES 195	466020	
M00054586A:F05 ES 195		1948.O21.gz43_250024
M00054586C:H02 ES 195	466020	
M00054586D:A03 ES 195	466020 160320	1948.021.gz43_250024

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CloneID	ES No	ClusterID	SequenceName
M00054586D:G01	ES 195	559419	1957.A09.gz43_250202
M00054587A:A08	ES 195	558413	1957.A13.gz43_250266
M00054587A;F06	ES 195	471181	1957.A16.gz43_250314
M00054587A:F09	ES 195	452506	1957.A18.gz43_250346
M00054587B:F06	ES 195	562174	1957.A23.gz43_250426
M00054587C:G02	ES 195	535129	1957.B03.gz43_250107
M00054587D:D04	ES 195	550322	1957.B05.gz43_250139
M00054588A:G08	ES 195	550855	1957.B11.gz43_250235
M00054588B:B12	ES 195	553331	1957.B13.gz43_250267
M00054588C:F04	ES 195	550217	1957.B19.gz43_250363
M00054589A;D09	ES 195	555920	1957.C04.gz43_250124
M00054589A:E05	ES 195	477097	1957.C05.gz43_250140
M00054589B:A07	ES 195	446164	1957.C06.gz43_250156
M00054589B:F05	ES 195	551463	1957.C10.gz43_250220
M00054589B:G10	ES 195	43656	1957.C13.gz43_250268
M00054589C:G11	ES 195	427723	1957.C18.gz43_250348
M00054589D:C06	ES 195	556376	1957.C20.gz43_250380
M00054590A:A01	ES 195	447311	1957.C23.gz43_250428
M00054590A:C10	ES 195	553755	1957.D02.gz43_250093
M00054590A:E02	ES 195	550594	1957.D03.gz43_250109
M00054590B:B11	ES 195	550088	1957.D06.gz43_250157
M00054590C:A03	ES 195	498827	1957.D08.gz43_250189
M00054590D:B01	ES 195	550084	1957.D16.gz43_250317
M00054590D:C12	ES 195	551975	1957.D18.gz43_250349
M00054591C:A01	ES 195	550044	1957.E04.gz43_250126
M00054591C:H09	ES 195	554764	1957.E06.gz43_250158
M00054591D:G11	ES 195	550886	1957.E11.gz43_250238
M00054592A:D09	ES 195	550315	1957.E17.gz43_250334
M00054592C:A05	ES 195	549964	1957.E20.gz43_250382
M00054593A:D01	ES 195	547509	1957.F03.gz43_250111
M00054593B:B03	ES 195	473859	1957.F09.gz43_250207
M00054593B:E03	ES 195	557970	1957.F12.gz43_250255
M00054593D:B11	ES 195	550153	1957.F18.gz43_250351
M00054594A:D04	ES 195	561836	1957.G01.gz43_250080
M00054594C:E06	ES 195	498777	1957.G08.gz43_250192
M00054595B:A11	ES 195	161489	1957.G20.gz43_250384
M00054595B:C12	ES 195	560357	1957.G23.gz43_250432
M00054595B:H09	ES 195	485431	1957.H06.gz43_250161
M00054595C:G06	ES 195	549665	1957.H12.gz43_250257
M00054595D:A10	ES 195	485029	1957.H18.gz43_250353
M00054596A:G11	ES 195	549731	1957.H24.gz43_250449
M00054596B:C01	ES 195	447356	1957.I04.gz43_250130
M00054596C:A06	ES 195	451594	1957.I11.gz43_250242
M00054596C:F09	ES 195	549609	1957.I14.gz43_250290

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M00054596C:F10	ES 195	455298	1957.I15.gz43_250306
M00054596C:G09	ES 195	561130	1957.I17.gz43_250338
M00054597A:A07	ES 195	467035	1957.I23.gz43_250434
M00054597A:C11	ES 195	558332	1957.J01.gz43_250083
M00054597A:D08	ES 195	558981	1957.J02.gz43_250099
M00054597A:G08	ES 195	549781	1957.J03.gz43_250115
M00054597B:F09	ES 195	493622	1957.J08.gz43_250195
M00054597D:E12	ES 195	461718	1957.J17.gz43_250339
M00054597D:F08	ES 195	449649	1957.J19.gz43_250371
M00054599B:A12	ES 195	446783	1957.K18.gz43_250356
M00054599B:C06	ES 195	549228	1957.K20.gz43_250388
M00054599B:F09	ES 195	549611	1957.K23.gz43_250436
M00054599C:E05	ES 195	561593	1957.L06.gz43_250165
M00054599D:B07	ES 195	559883	1957.L08.gz43_250197
M00054600A:G12	ES 195	559554	1957.L17.gz43_250341
M00054600A:H03	ES 195	549816	1957.L18.gz43_250357
M00054600C:B10	ES 195	459581	1957.L20.gz43_250389
M00054600C:D03	ES 195	554193	1957.L22.gz43_250421
M00054600D:H07	ES 195	554869	1957.M08.gz43 250198
M00054601A:E08	ES 195	555949	1957.M09.gz43_250214
M00054601B:D08	ES 195	446503	1957.M12.gz43 250262
M00054602A:B03	ES 195	550085	1957.M23.gz43_250438
M00054602A:E06	ES 195	416884	1957.N05.gz43_250151
M00054602C:B06	ES 195	547509	1957.N15.gz43_250311
M00054602C:C12	ES 195	487176	1957.N16.gz43_250327
M00054602D:A06	ES 195	562722	1957.N20.gz43_250391
M00054603A:G06	ES 195	552907	1957.O07.gz43_250184
M00054603A:G12	ES 195	552913	1957.O09.gz43_250216
M00054603B:B04	ES 195	526733	1957.O11.gz43_250248
M00054603B:C09	ES 195	447308	1957.O13.gz43 250280
M00054603B:G08	ES 195	356058	1957.O16.gz43_250328
M00054603C:G09	ES 195	408130	1957.O18.gz43_250360
M00054603D:B02	ES 195	552121	1957.O20.gz43_250392
M00054604C:B05	ES 195	562529	1957.P07.gz43 250185
M00054604C:F03	ES 195	549599	1957.P12.gz43 250265
M00054604D:A04	ES 195	563514	1957.P15.gz43 250313
M00054605A:A03	ES 195	558927	1957.P18.gz43 250361
M00054605B:F10	ES 195	460493	1957.P24.gz43 250457
M00054605B:G01	ES 195	549649	1958.A02.gz43_250474
M00054605C:A04	ES 195	548864	1958.A05.gz43_250522
M00054605C:D03	ES 195	188753	1958.A06.gz43_250538
M00054605C:H02	ES 195	549810	1958.A09.gz43_250586
M00054606A:A03	ES 195	449994	1958.A13.gz43_250650
M00054606A:D02	ES 195	562508	1958.A16.gz43 250698

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00054606C:B06	ES 195	549131	1958.A23.gz43_250810
M00054606C:E04	ES 195	549481	1958.B01.gz43_250459
M00054606D:B05	ES 195	557190	1958.B02.gz43_250475
M00054607A:B06	ES 195	560862	1958.B05.gz43_250523
M00054607A:G02	ES 195	552823	1958.B07.gz43_250555
M00054608B:D08	ES 195	490393	1958.B19.gz43_250747
M00054608C:H04	ES 195	549889	1958.B24.gz43_250827
M00054609A:D01	ES 195	423947	1958.C05.gz43_250524
M00054609A:F01	ES 195	561892	1958.C08.gz43_250572
M00054609A:H04	ES 195	549912	1958.C09.gz43_250588
M00054609B:E01	ES 195	556530	1958.C12.gz43_250636
M00054609B:H11	ES 195	507349	1958.C13.gz43_250652
M00054609C:G06	ES 195	549779	1958.C17.gz43_250716
M00054609D:E12	ES 195	490152	1958.C22.gz43_250796
M00054609D:H06	ES 195	549829	1958.C24.gz43 250828
M00054610A:E05	ES 195	558332	1958.D05.gz43 250525
M00054611B:F12	ES 195	549626	1958.E13.gz43 250654
M00054611B:G09	ES 195	478511	1958.E14.gz43_250670
M00054611C:C02	ES 195	561144	1958.E16.gz43 250702
M00054611C:E01	ES 195	555660	1958.E18.gz43_250734
M00054612A:D12	ES 195	549347	1958.F08.gz43 250575
M00054612B:B11	ES 195	549114	1958.F10.gz43 250607
M00054612B:E02	ES 195	488108	1958.F12.gz43 250639
M00054612D:B12	ES 195	552364	1958.F22.gz43 250799
M00054613A:D07	ES 195	550177	1958.G06.gz43 250544
M00054613B:H04	ES 195	550018	1958.G11.gz43 250624
M00054613D:H09	ES 195	553028	1958.G21.gz43_250784
M00054614A:G11	ES 196	557559	1958.G24.gz43 250832
M00054614B:E01	ES 196	405042	1958.H01.gz43_250465
M00054615B:E03	ES 196	497477	1958.H16.gz43_250705
M00054615C:D09	ES 196	557947	1958,H23,gz43 250817
M00054616A:H01	ES 196	553758	1958.I15.gz43 250690
M00054616A:H03	ES 196	481360	1958.I16.gz43 250706
M00054616D:C10	ES 196	552418	1958,J01,gz43 250467
M00054616D:G09	ES 196	549739	1958.J03.gz43 250499
M00054616D:H06	ES 196	513115	1958.J05.gz43 250531
M00054617A:C07	ES 196	553453	1958,J08,gz43 250579
M00054617B:A09	ES 196	548965	1958.J10.gz43 250611
M00054617B:C05	ES 196	549233	1958.J12.gz43_250643
M00054617B:D06	ES 196	549388	1958,J13.gz43 250659
M00054617B:B04	ES 196	549122	1958.K06.gz43_250548
M00054618C:H02	ES 196	555371	1958.K14.gz43 250676
M00054618D:D04	ES 196	552437	1958.K14.gz43_250070
TATOROD-101 OD.DO-	120 120	332437	1730.1510.BZ43_Z30700

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CloneID	ES No	ClusterID	SequenceName
M00054618D:E06	ES 196	490890	1958.K19.gz43_250756
M00054618D:E11	ES 196	451009	1958.K20.gz43_250772
M00054619A:C05	ES 196	447913	1958.K24.gz43_250836
M00054619C:F04	ES 196	554036	1958.L09.gz43_250597
M00054619D:C10	ES 196	562749	1958.L12.gz43_250645
M00054620A:C09	ES 196	553204	1958.L15.gz43_250693
M00054620B:B10	ES 196	552194	1958,L20,gz43_250773
M00054620C:C08	ES 196	448276	1958.L22.gz43_250805
M00054620C:C12	ES 196	560491	1958.L23.gz43_250821
M00054620D:D06	ES 196	561116	1958.M03.gz43_250502
M00054620D:D11	ES 196	473111	1958.M04.gz43_250518
M00054620D:F11	ES 196	556896	1958.M06.gz43_250550
M00054620D:G11	ES 196	395411	1958.M07.gz43_250566
M00054621B:A10	ES 196	548859	1958.M13.gz43_250662
M00054621C:G03	ES 196	451025	1958.M20.gz43_250774
M00054621D:A10	ES 196	518049	1958.M24.gz43_250838
M00054621D:C06	ES 196	558642	1958.N01.gz43_250471
M00054621D:D11	ES 196	549320	1958.N02.gz43_250487
M00054622A:D06	ES 196	27480	1958.N07.gz43_250567
M00054622A:H01	ES 196	561068	1958.N08.gz43_250583
M00054622B:F05	ES 196	491644	1958.N10.gz43_250615
M00054622B:H09	ES 196	556308	1958.N12.gz43_250647
M00054622D:C02	ES 196	562932	1958.N18.gz43_250743
M00054622D:D10	ES 196	551463	1958.N20.gz43_250775
M00054622D:F05	ES 196	485504	1958.N21.gz43_250791
M00054622D:G11	ES 196	556835	1958.N22.gz43_250807
M00054622D:H03	ES 196	453864	1958.N23.gz43 250823
M00054623B:B11	ES 196	556424	1958.O06.gz43_250552
M00054623C:E09	ES 196	558175	1958.009.gz43_250600
M00054623D:C12	ES 196	551305	1958.O12.gz43_250648
M00054624A:B11	ES 196	550164	1958.O16.gz43_250712
M00054624B:G10	ES 196	559938	1958.O21.gz43_250792
M00054624D:B06	ES 196	554810	1958.O23.gz43_250824
M00054625A:D07	ES 196	558103	1958.P04.gz43 250521
M00054625A:E05	ES 196	557199	1958.P05.gz43_250537
M00054625B:B02	ES 196	551250	1958.P07.gz43 250569
M00054625D:F06	ES 196	562712	1958.P21.gz43 250793
M00054625D:H07	ES 196	461734	1958.P22.gz43_250809
M00054627A:B08	ES 196	556790	1959.A20.gz43_254083
M00054628C:G11	ES 196	557559	1959.C14.gz43_253989
M00054629C:G06	ES 196	555923	1959.D08.gz43 253894
M00054630A:D08	ES 196	562541	1959.D17.gz43_254038
M00054630B:A06	ES 196	551975	1959.D19.gz43_254070
M00054631D:C02	ES 196	552092	1959.F03.gz43_253816

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CloneID	ES No	ChisterID	SequenceName
M00054635B:G05	ES 196	268197	1959.H16.gz43_254026
M00054638D:F02	ES 196	555949	1959.K24.gz43_254157
M00054640B:C05	ES 196	549210	1959.M04.gz43_253839
M00054642D:F11	ES 196	549516	1959.N22.gz43_254128
M00054643B:G09	ES 196	552618	1959.O10.gz43 253937
M00054644B:F02	ES 196	557345	1959.P08.gz43_253906
M00054644D:F11	ES 196	549624	1959.P20.gz43_254098
M00054647D:H02	ES 196	502683	1960.B23.gz43_254516
M00054648C:H10	ES 196	450867	1960.C12.gz43_254341
M00054650A:B08	ES 196	410667	1960.D12.gz43_254342
M00054651D:D02	ES 196	452662	1960.E24.gz43_254535
M00054652B:A04	ES 196	387728	1960.F04.gz43_254216
M00054656C:E10	ES 196	561590	1960.I07.gz43_254267
M00054657D:E05	ES 196	551531	1960.J11.gz43_254332
M00054661A:B03	ES 196	193991	1960.L03.gz43_254206
M00054661D:A07	ES 196	551065	1960.L12.gz43_254350
M00054663D:D09	ES 196	466851	1960.M24.gz43_254543
M00054664D:A01	ES 196	264575	1960.N14.gz43_254384
M00054668A:D01	ES 196	450624	1969.A10.gz43_254691
M00054670D:F10	ES 196	556424	1969.B21.gz43_254868
M00054671A:H07	ES 196	579481	1969.C07.gz43_254645
M00054671D:A12	ES 196	573733	1969.C24.gz43_254917
M00054672C:F01	ES 196	140763	1969.D20.gz43_254854
M00054673B:G08	ES 196	551845	1969.E09.gz43_254679
M00054674B:B03	ES 196	551269	1969.F03.gz43_254584
M00054677D:H08	ES 196	556959	1969.I02.gz43_254571
M00054683D:F01	ES 196	551630	1969,M04.gz43_254607
M00054685D:D09	ES 196	415326	1969.N11.gz43_254720
M00054689C:B11	ES 196	557935	1970.A03.gz43_263422
M00054689D:E12	ES 196	551527	1970.A06.gz43_263470
M00054690B:B02	ES 196	559389	1970.A14.gz43_263598
M00054690B:D10	ES 196	552418	1970.A16.gz43_263630
M00054690D:G03	ES 196	549388	1970.B04.gz43_263439
M00054692B:C06	ES 196	550330	1970.B20.gz43_263695
M00054692B:D01	ES 196	420686	1970.B21.gz43_263711
M00054692C:B02	ES 196	42994	1970.B23.gz43_263743
M00054692D:F09	ES 196	479208	1970.B24.gz43_263759
M00054693A:C09	ES 196	551243	1970.C05.gz43_263456
M00054693A:E06	ES 196	558938	1970.C08.gz43_263504
M00054693B:B01	ES 196	559036	1970.C09.gz43_263520
M00054693C:A02	ES 196	549995	1970.C14.gz43_263600
M00054693D:A05	ES 196	551150	1970.C18.gz43_263664
M00054693D:A08	ES 196	. 448332	1970.C19.gz43_263680
M00054693D:C04	ES 196	450755	1970.C20.gz43_263696

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CloneID	ES No	ClusterID	SequenceName
M00054695A:A12	ES 196	552051	1970.D08.gz43_263505
M00054696B:H11	ES 196	553128	1970.E06.gz43_263474
M00054697A:E03	ES 196	552649	1970.E12.gz43_263570
M00054697A:G06	ES 196	562216	1970.E14.gz43_263602
M00054697C:E11	ES 196	552581	1970,E21.gz43_263714
M00054699A:G01	ES 196	584499	1970.F15.gz43_263619
M00054699B:A05	ES 196	551996	1970.F16.gz43_263635
M00054699B:F05	ES 196	552733	1970.F18.gz43_263667
M00054699C:A04	ES 196	551995	1970.F21.gz43_263715
M00054699D:A12	ES 196	552055	1970.G02.gz43_263412
M00054700A:B03	ES 196	552254	1970.G04.gz43_263444
M00054700A:F12	ES 196	552733	1970.G06.gz43_263476
M00054701D:H05	ES 196	555255	1970.G23.gz43_263748
M00054702A;C06	ES 196	556896	1970.H01.gz43_263397
M00054702A:F03	ES 196	483141	1970.H02.gz43_263413
M00054702A:H03	ES 196	509202	1970,H03.gz43_263429
M00054702C:G06	ES 196	551754	1970.H09.gz43_263525
M00054702D:E07	ES 196	389201	1970.H12.gz43_263573
M00054702D:F05	ES 196	452392	1970.H14.gz43_263605
M00054702D:H10	ES 196	549945	1970.H16.gz43_263637
M00054703C:F01	ES 196	521552	1970.H22.gz43_263733
M00054703D:E07	ES 196	549238	1970.I01.gz43_263398
M00054703D:F04	ES 196	561434	1970.I02.gz43_263414
M00054704B:B11	ES 196	549038	1970.I06.gz43_263478
M00054704D:F02	ES 196	552682	1970.I12.gz43_263574
M00054705B:C08	ES 196	552314	1970.I14.gz43_263606
M00054705B:D02	ES 196	527679	1970.I16.gz43_263638
M00054705B:E04	ES 196	498454	1970.I18.gz43_263670
M00054705C:B02	ES 196	475730	1970.I21.gz43_263718
M00054705C:D11	ES 196	554477	1970.I24.gz43_263766
M00054705D:G03	ES 196	184995	1970.J03.gz43_263431
M00054706A;G10	ES 196	491827	. 1970.J09.gz43_263527
M00054706B:F10	ES 196	562884	1970.J15.gz43_263623
M00054706C:A04	ES 196	551955	1970.J17.gz43_263655
M00054706C:B12	ES 196	550053	1970.J19.gz43_263687
M00054706D:B03	ES 196	553358	1970.J22.gz43_263735
M00054707B:B08	ES 196	453708	1970.K02.gz43_263416
M00054707C:D02	ES 196	549151	1970.K06.gz43_263480
M00054707C:G07	ES 196	552894	1970.K09.gz43_263528
M00054708A:F11	ES 196	552686	1970.K12.gz43_263576
M00054708B:F04	ES 196	391511	1970.K16.gz43_263640
M00054709A:A10	ES 196	560317	1970.K24.gz43_263768
M00054710B:B10	ES 196	552197	1970.L17.gz43_263657
M00054710C:A12	ES 196	558768	1970.L22.gz43_263737

Table 13

Table 15			
CloneID	ES No	ClusterID	SequenceName
M00054710D:A02	ES 196	551941	1970.M01.gz43_263402
M00054711A:B05	ES 196	553685	1970.M05.gz43_263466
M00054711C:A08	ES 196	552073	1970.M11.gz43_263562
M00054712C:A04	ES 196	558046	1970.N01.gz43_263403
M00054712C:C07	ES 196	549557	1970.N05.gz43_263467
M00054713B:B10	ES 196	551479	1970.N14.gz43_263611
M00054713C:D07	ES 196	553938	1970.N18.gz43_263675
M00054714A:C05	ES 196	430917	1970.O01.gz43_263404
M00054714B:F05	ES 196	32021	1970.O06.gz43_263484
M00054714C:E01	ES 196	552625	1970.O09.gz43_263532
M00054715A:C06	ES 196	452559	1970.O15.gz43_263628
M00054715A:G02	ES 196	550975	1970.O17.gz43_263660
M00054715C:D05	ES 196	451032	1970.O23.gz43_263756
M00054715C:D11	ES 196	419281	1970.O24.gz43_263772
M00054715D:E11	ES 196	552598	1970.P07.gz43_263501
M00054715D:F09	ES 196	583641	1970.P08.gz43_263517
M00054716B:D06	ES 196	490393	1970.P16.gz43_263645
M00054716D:D03	ES 196	552497	1970.P23.gz43_263757
M00054717A:A03	ES 196	552086	1971.A01.gz43_246992
M00054717A:C07	ES 196	549550	1971.A03.gz43_247024
M00054717A:G09	ES 196	450278	1971.A06.gz43_247072
M00054717C:G07	ES 196	554913	1971.A18.gz43_247264
M00054718A:D11	ES 196	555571	1971.A24.gz43_247360
M00054718B:D03	ES 196	555837	1971.B05.gz43_247057
M00054719A:A02	ES 196	552028	1971.B13.gz43_247185
M00054719A:H06	ES 196	553012	1971.B16.gz43_247233
M00054720C:A01	ES 196	553123	1971.D01.gz43_246995
M00054720C:G10	ES 196	557714	1971.D05.gz43_247059
M00054720C:H01	ES 196	561313	1971.D06.gz43_247075
M00054721C:D11	ES 196	446999	1971.E05.gz43_247060
M00054722C:D01	ES 196	450410	1971.F05.gz43_247061
M00054722D:C08	ES 196	553316	1971.F09.gz43_247125
M00054724A:G02	ES 196	421959	1971.G14.gz43_247206
M00054724B:F06	ES 196	551826	1971.G18.gz43_247270
M00054724D:C05	ES 196	561636	1971.H01.gz43_246999
M00054724D:G12	ES 196	553805	1971.H07.gz43_247095
M00054725A:A03	ES 196	448136	1971.H09.gz43_247127
M00054725A:A05	ES 196	551096	1971.H10.gz43_247143
M00054725A:F09	ES 196	555491	1971.H11.gz43_247159
M00054725A:H06	ES 196	86175	1971.H12.gz43_247175
M00054725B:G12	ES 196	555676	1971.H19.gz43_247287
M00054725D:F04	ES 196	553731	1971.I04.gz43_247048
M00054726A:B11	ES 196	552933	1971.I07.gz43_247096
M00054726A:D02	ES 196	555818	1971.I09.gz43_247128

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00054726A:E01	ES 197	551487	1971.I10.gz43_247144
M00054726B:F08	ES 197	553739	1971.I16.gz43_247240
M00054726B:F09	ES 197	557747	1971.I17.gz43_247256
M00054726C:D10	ES 197	449035	1971.I21.gz43_247320
M00054726C:E07	ES 197	524721	1971.I22.gz43_247336
M00054727A:B12	ES 197	553264	1971.J09.gz43_247129
M00054727D:C06	ES 197	496772	1971.K01.gz43_247002
M00054728B:C10	ES 197	557356	1971.K09.gz43_247130
M00054728C:B08	ES 197	559113	1971.K13.gz43_247194
M00054728D:B07	ES 197	554048	1971.K18.gz43_247274
M00054729B:A08	ES 197	484043	1971.L05.gz43_247067
M00054729B:B03	ES 197	557426	1971.L06.gz43_247083
M00054729B:E06	ES 197	451812	1971.L09.gz43_247131
M00054729C:C08	ES 197	561830	1971.L12.gz43_247179
M00054729C:G07	ES 197	562263	1971.L15.gz43_247227
M00054729D:D05	ES 197	550315	1971.L18.gz43_247275
M00054730B:F11	ES 197	559380	1971.M11.gz43_247164
M00054731A;D07	ES 197	452488	1971.N05.gz43_247069
M00054731B:C12	ES 197	553310	1971.N13.gz43_247197
M00054731B:D04	ES 197	448453	1971.N14.gz43_247213
M00054731B:G02	ES 197	553751	1971.N17.gz43_247261
M00054731C:H12	ES 197	553869	1971.N24.gz43_247373
M0005473 1D:H11	ES 197	215005	1971.004.gz43_247054
M00054732A:B07	ES 197	561379	1971.006.gz43_247086
M00054734B:C06	ES 197	556654	1971.P10.gz43_247151
M00054734B:G10	ES 197	549288	1971.P15.gz43_247231
M00054734C:A07	ES 197	553108	1971.P17.gz43_247263
M00054735A:G09	ES 197	552686	1971.P24.gz43_247375
M00054736B:H03	ES 197	522322	1972.B13.gz43_247569
M00054737B:H11	ES 197	556167	1972.C05.gz43_247442
M00054737C:B01	ES 197	454186	1972.C06.gz43_247458
M00054738B:E12	ES 197	560282	1972.C21.gz43_247698
M00054738D:F01	ES 197	558679	1972.D13.gz43_247571
M00054738D:G07	ES 197	495832	1972.D15.gz43_247603
M00054739A:F07	ES 197	555773	1972.D24.gz43_247747
M00054739C:B12	ES 197	466235	1972.E08.gz43_247492
M00054739C:E05	ES 197	450142	1972.E12.gz43_247556
M00054740C:H08	ES 197	493122	1972.F15.gz43_247605
M00054741C:D08	ES 197	493135	1972.G15.gz43_247606
M00054741D:C05	ES 197	551778	1972.G19.gz43_247670
M00054743A:C07	ES 197	424723	1972.I03.gz43_247416
M00054743A:E04	ES 197	473455	1972.I04.gz43_247432
M00054743C:E11	ES 197	452257	1972.I13.gz43_247576

Table 13

CloneID	ES No	ClusterID	SequenceName
M00054743D:F04	ES 197	555077	1972.I19.gz43_247672
M00054744A:G10	ES 197	553834	1972.J05.gz43_247449
M00054744C:D02	ES 197	587854	1972.J16.gz43_247625
M00054744D:G02	ES 197	557068	1972.K03.gz43_247418
M00054745C:B12	ES 197	562793	1972.K15.gz43_247610
M00054745C:H02	ES 197	557871	1972.K19.gz43 247674
M00054745D:G09	ES 197	551441	1972.L04.gz43_247435
M00054746A:H02	ES 197	559494	1972.L09.gz43_247515
M00054746D:E05	ES 197	503452	1972.L22.gz43_247723
M00054747D:B05	ES 197	474309	1972.M24.gz43_247756
M00054748B:G10	ES 197	523753	1972.N13.gz43_247581
M00054748B:H09	ES 197	59202	1972.N14.gz43_247597
M00054750A:E11	ES 197	446997	1972.O06.gz43_247470
M00054750A;G10	ES 197	562323	1972.O09.gz43_247518
M00054750B:F02	ES 197	557568	1972.O12.gz43_247566
M00054750C:D01	ES 197	553457	1972.O18.gz43_247662
M00054750D:F01	ES 197	559215	1972.P01.gz43_247391
M00054750D:H12	ES 197	499517	1972.P03.gz43_247423
M00054751A:A05	ES 197	86145	1972.P04.gz43_247439
M00054751B:F12	ES 197	553702	1972.P14.gz43_247599
M00054751D:G10	ES 197	553800	1972.P24.gz43_247759
M00054752A:C12	ES 197	562451	1981.A03.gz43_247792
M00054752B:H06	ES 197	495942	1981.A10.gz43_247904
M00054753A:A05	ES 197	554000	1981.A18.gz43_248032
M00054753C:H02	ES 197	553918	1981.B08.gz43_247873
M00054753D:A03	ES 197	477046	1981.B09.gz43_247889
M00054753D:C12	ES 197	256179	1981.B11.gz43_247921
M00054753D:H10	ES 197	553922	1981.B13.gz43_247953
M00054754B:F04	ES 197	558609	1981.B16.gz43_248001
M00054754C:H09	ES 197	550580	1981.B22.gz43_248097
M00054754D:F11	ES 197	551693	1981.C04.gz43_247810
M00054755A:E10	ES 197	493135	1981.C10.gz43_247906
M00054755C:D04	ES 197	561412	1981.C24.gz43_248130
M00054755D:E02	ES 197	562360	1981.D02.gz43_247779
M00054755D:E05	ES 197	513632	1981.D03.gz43_247795
M00054756A:C12	ES 197	551289	1981.D09.gz43_247891
M00054756C:C08	ES 197	551288	1981.D15.gz43_247987
M00054756D:F05	ES 197	451993	1981.D19.gz43_248051
M00054757A:H07	ES 197	555343	1981.D21.gz43_248083
M00054757B:H04	ES 197	561433	1981.E04.gz43_247812
M00054758A;F03	ES 197	554336	1981.E13.gz43_247956
M00054758B:C10	ES 197	122169	1981.E15.gz43_247988
M00054758B:D03	ES 197	554352	1981.E16.gz43_248004
M00054758B:H03	ES 197	554049	1981.E18.gz43_248036

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Table 13

CloneID	ES No	ClusterID	SequenceNam
M00054758D:D03	ES 197	551415	1981.E24.gz43_24813
M00054759A:B08	ES 197	553158	1981.F03.gz43_24779
M00054759C:D07	ES 197	553800	1981.F11.gz43_24792
M00054759C:G10	ES 197	560529	1981.F13.gz43_24795
M00054759D:E01	ES 197	556064	1981.F17.gz43_24802
M00054760A;A12	ES 197	551117	1981.F22.gz43_24810
M00054760A:D10	ES 197	561535	1981.F24.gz43_24813
M00054760B:A10	ES 197	549588	1981.G04.gz43_24781
M00054760D:B03	ES 197	263800	1981.G09.gz43_24789
M00054761C:E02	ES 197	558719	1981.G24.gz43_24813
M00054761D:C11	ES 197	493604	1981.H04,gz43 24781
M00054762A:D09	ES 197	556183	1981.H10.gz43 24791
M00054762B:F02	ES 197	558900	1981.H14.gz43 24797
M00054762B:F07	ES 197	551617	1981.H15.gz43_24799
M00054762C:A12	ES 197	558503	1981.H18.gz43_24803
M00054762D:C06	ES 197	551338	1981.H24.gz43 24813
M00054763A:A10	ES 197	472196	1981.I02.gz43 24778
M00054763C:D07	ES 197	554021	1981.I04.gz43 2478
M00054763C:F10	ES 197	558900	. 1981.I05.gz43_24783
M00054763C:H04	ES 197	450840	1981.I06.gz43 2478-
M00054764A:E11	ES 197	576803	1981.I14.gz43 24797
M00054764C:G04	ES 197	555754	1981.I22.gz43 24810
M00054764D:F01	ES 197	556019	1981.J01.gz43 2477
M00054765B:C03	ES 197	561487	1981.J07.gz43_2478
M00054765B:C11	ES 197	554181	1981.J09.gz43 24789
M00054765C:F10	ES 197	553131	1981.J20.gz43 2480'
M00054765D:D05	ES 197	554294	1981.K05.gz43 2478
M00054766A:H10	ES 197	508126	1981.K17.gz43 24802
M00054766C:B08	ES 197	498662	1981.L03.gz43 24780
M00054766C:E01	ES 197	552541	1981.L07.gz43 2478
M00054766D:H02	ES 197	260558	1981.L15,gz43_2479
M00054766D:H12	ES 197	554851	1981.L16.gz43_2480
M00054767A:F08	ES 197	556475	1981.L22.gz43 2481
M00054767C:C08	ES 197	552535	1981.M13.gz43_2479
M00054767C:D03	ES 197	455220	1981.M14.gz43_2479
M00054767C:H06	ES 197	549576	1981.M17.gz43 2480
M00054767D;G09	ES 197	552549	1981.M22.gz43 2481
M00054768B:B05	ES 197	559369	1981.N01.gz43 2477
M00054768B:D10	ES 197	400628	1981.N05.gz43 2478
M00054768D:A01	ES 197	448357	1981.N10.gz43_2479
M00054768D:B11	ES 197	451051	1981.N11.gz43 2479
M00054769A:G10	ES 197	512432	1981.N21.gz43 2480
M00054769B:D12	ES 197	550402	1981.O04.gz43 24782
M00054769B:F03	ES 197	554635	1981.O06.gz43_2478:

Table 13

Table 13			
CloneID	ES No	ClusterID	SequenceName
M00054770A:C06	ES 197	558949	1981.O19.gz43_248062
M00054770B:A12	ES 197	552733	1981.O24.gz43_248142
M00054770B:B11	ES 197	556918	1981.P01.gz43_247775
M00054770B:D09	ES 197	448510	1981.P04.gz43 247823
M00054770C:A04	ES 197	528775	1981.P07.gz43_247871
M00054770C:C04	ES 197	554161	1981.P10.gz43_247919
M00054770C:D05	ES 197	555478	1981.P11.gz43_247935
M00054770C:F10	ES 197	554581	1981,P12.gz43_247951
M00054771A:E01	ES 197	470351	1981,P23,gz43_248127
M00054771B:F12	ES 197	492779	1982.A02.gz43_248160
M00054771C:A11	ES 197	551068	1982.A04.gz43_248192
M00054771D:G01	ES 197	551718	1982.A08.gz43_248256
M00054772B:B01	ES 197	586600	1982.A15.gz43_248368
M00054772B:D04	ES 197	472672	1982.A17.gz43_248400
M00054772B:F03	ES 197	493085	1982.A18.gz43_248416
M00054772D:G12	ES 197	553774	1982.B06.gz43_248225
M00054773A:E09	ES 197	551169	1982.B08.gz43_248257
M00054773A:H05	ES 197	402147	1982.B09.gz43_248273
M00054773C:D06	ES 197	42323	1982.B15.gz43_248369
M00054773C:F09	ES 197	557834	1982.B18.gz43_248417
M00054773C:F10	ES 197	448629	1982.B19.gz43_248433
M00054774B:A07	ES 197	555686	1982.C03.gz43_248178
M00054774B:G08	ES 197	554072	1982.C06.gz43_248226
M00054774C:A03	ES 197	552019	1982.C07.gz43_248242
M00054774C:D12	ES 197	552495	1982.C08.gz43_248258
M00054774D:D06	ES 197	559050	1982,C10.gz43_248290
M00054775A:D07	ES 197	555462	1982.C16.gz43_248386
M00054775A:G03	ES 197	595181	1982.C17.gz43_248402
M00054776A:D10	ES 197	557269	1982.D09.gz43_248275
M00054776A:F01	ES 197	560520	1982.D10.gz43_248291
M00054776A:F07	ES 197	447926	1982.D11.gz43_248307
M00054776A:G01	ES 197	412621	1982.D12.gz43_248323
M00054776C:G06	ES 197	554828	1982.E06.gz43_248228
M00054776D:G09	ES 197	553743	1982.E11.gz43_248308
M00054778A:D01	ES 197	555958	1982,F01.gz43_248149
M00054778A:F08	ES 197	551003	1982.F05.gz43_248213
M00054778B:A02	ES 197	551080	1982.F08.gz43_248261
M00054778C:F09	ES 197	548998	1982.F18.gz43_248421
M00054779B:A07	ES 197	465207	1982.G05.gz43_248214
M00054779B:B07	ES 197	559872	1982.G06.gz43_248230
M00054779B:B11	ES 197	556183	1982.G07.gz43_248246
M00054779D:F07	ES 197	554627	1982.G17.gz43_248406
M00054779D:F08	ES 197	551425	1982.G18.gz43_248422
M00054780A:B06	ES:197	550422	1982.G21.gz43_248470

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CloneID	ES No	ClusterID	SequenceName
M00054780B:B06	ES 197	555524	1982.H03.gz43_248183
M00054780B:E03	ES 197	533991	1982.H05.gz43_248215
M00054780B:G10	ES 197	491570	1982.H06.gz43_248231
M00054780C:G11	ES 197	558519	1982.H14.gz43_248359
M00054781A:H09	ES 197	505792	1982.H21.gz43_248471
M00054781D:F10	ES 197	556171	1982.I02.gz43_248168
M00054782B:B02	ES 197	551167	1982.I05.gz43_248216
M00054782B:D09	ES 197	551425	1982.I07.gz43_248248
M00054782D:D12	ES 197	575924	1982.I10.gz43_248296
M00054783C:C02	ES 197	557039	1982.I16.gz43_248392
M00054783C:C03	ES 197	552325	1982.I17.gz43_248408
M00054784C:D11	ES 197	552535	1982.J07.gz43_248249
M00054784D:B01	ES 197	554582	1982.J10.gz43_248297
M00054785B:B07	ES 197	552188	1982.J13.gz43_248345
M00054785C:C02	ES 197	562263	1982.J15.gz43_248377
M00054785C:G11	ES 197	462511	1982.J17.gz43_248409
M00054785D:A07	ES 197	552005	1982.J19.gz43_248441
M00054786A:G11	ES 198	561422	1982.J21.gz43_248473
M00054786C:G01	ES 198	553330	1982.K02.gz43_248170
M00054786D:F08	ES 198	553868	1982.K05.gz43_248218
M00054787D:A10	ES 198	580179	1982.K12.gz43_248330
M00054788C:G04	ES 198	552813	1982.K17.gz43_248410
M00054789A:A02	ES 198	490414	1982.K18.gz43_248426
M00054790D:D05	ES 198	548874	1982.L06.gz43_248235
M00054801C:G01	ES 198	549233	1982.L17.gz43_248411
M00054802A:G03	ES 198	446394	1982.M01.gz43_248156
M00054802A:H05	ES 198	490507	1982.M03.gz43_248188
M00054802C:A07	ES 198	374282	1982.M06.gz43_248236
M00054802D:A09	ES 198	555696	1982.M10.gz43_248300
M00054802D:C02	ES 198	497477	1982.M12.gz43_248332
M00054802D:C03	ES 198	456052	1982.M13.gz43_248348
M00054803A:D08	ES 198	175758	1982.M17.gz43_248412
M00054803A:E10	ES 198	551553	1982.M18.gz43_248428
M00054803B:B12	ES 198	554116	1982,M20,gz43_248460
M00054803C:G01	ES 198	551755	1982,M24.gz43_248524
M00054804A:G08	ES 198	496570	1982.N08.gz43_248269
M00054804A:H04	ES 198	539353	1982.N09.gz43_248285
M00054804B:E07	ES 198	553979	1982.N11.gz43_248317
M00054804C:F04	ES 198	553237	1982.N14.gz43_248365
M00054804D:A11	ES 198	559896	1982.N16.gz43_248397
M00054804D:D07	ES 198	555571	1982.N18.gz43_248429
M00054804D:H12	ES 198	417259	1982.N22.gz43_248493
M00054805A:E09	ES 198	553612	1982.O03.gz43_248190

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1 abie 15			
CloneID	ES No	ClusterID	SequenceName
M00054805A:H04	ES 198	552183	1982.O04.gz43_248206
M00054805B:E06	ES 198	556856	1982.O10.gz43_248302
M00054805B:E11	ES 198	497912	1982.O11.gz43 248318
M00054805B:G02	ES 198	556856	1982.O14.gz43 248366
M00054805B:G10	ES 198	394168	1982,O16.gz43 248398
M00054806A:G04	ES 198	553848	1982.P02.gz43 248175
M00054806C:C12	ES 198	517146	1982.P08.gz43 248271
M00054806D:C01	ES 198	553331	1982.P10.gz43 248303
M00054811A;C10	ES 198	551512	1983.C10.gz43 255077
M00054811B:H12	ES 198	494471	1983.C19.gz43 255221
M00054811D:F01	ES 198	551659	1983.D01.gz43 254934
M00054812D:H05	ES 198	549816	1983.E06.gz43 255015
M00054813B:D11	ES 198	456520	1983.E19.gz43 255223
M00054814A:F07	ES 198	557525	1983.F09.gz43 255064
M00054823B:E07	ES 198	450788	1983.K11.gz43 255101
M00054824A:E02	ES 198	555641	1983,L07,gz43 255038
M00054826B:G08	ES 198	447087	1983.N07.gz43_255040
M00054831A:E11	ES 198	554196	1984.A12.gz43 255491
M00054831B:E05	ES 198	451081	1984.A20.gz43 255619
M00054839D:F08	ES 198	562302	1984.G07.gz43_255417
M00054840D:B03	ES 198	557703	1984.G21.gz43 255641
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Table 13

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Table 13

Table 15			
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M00054900C:C03	ES 198	516691	1995.H12.gz43_256266
M00054900C:D08	ES 198	556753	1995.H13.gz43_256282
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M00054901B:H10	ES 198	556165	1995.I03.gz43_256123
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Table 15			
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M00054904D:G04	ES 199	558868	1995.K20.gz43_256397
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M00054910D:F04	ES 199	555786	1995.N13.gz43_256288
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M00054925B:B01	ES 199	559728	1996.A04.gz43_256539
M00054927C:C03	ES 199	473512	1996.B13.gz43_256684
M00054928A;C11	ES 199	550782	1996.B20.gz43_256796
M00054930D:D04	ES 199	558024	1996.D13.gz43_256686
M00054931C:F10	ES 199	549171	1996.E06.gz43_256575
M00054933A:H09	ES 199	44608	1996.F18.gz43_256768
M00054933B:C02	ES 199	562106	1996.F20.gz43_256800
M00054936B:G02	ES 199	555739	1996.I18.gz43_256771
M00054936C:B08	ES 199	549805	1996.119.gz43_256787

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Table 15			
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M00054936C:G03	ES 199	447585	1996,I21.gz43_256819
M00054936D:H10	ES 199	554845	1996.J01.gz43_256500
M00054938A:B08	ES 199	418512	1996.K02.gz43_256517
M00054939A:B11	ES 199	560088	1996.L08.gz43_256614
M00054939D:E03	ES 199	552566	1996.M08.gz43_256615
M00054940C:H12	ES 199	227777	1996.N13.gz43_256696
M00054940D:E06	ES 199	553535	1996.N16.gz43_256744
M00054941A:A08	ES 199	558397	1996.N18.gz43_256776
M00054941C:G04	ES 199	562550	1996.O06.gz43_256585
M00054942A:G01	ES 199	549622	1996.O20.gz43_256809
M00054942C:B04	ES 199	67060	1996.P05.gz43_256570
M00054943A:H11	ES 199	481166	1996.P20.gz43_256810
M00054943D:F12	ES 199	562053	2005.A14.gz43_257085
M00054945B:G09	ES 199	554833	2005.C07.gz43_256975
M00054949C:H07	ES 199	430194	2005.F21.gz43_257202
M00054951A:G03	ES 199	555996	2005.H14.gz43_257092
M00054952A:B12	ES 199	496870	2005.I10.gz43_257029
M00054952D:A11	ES 199	555018	2005.J08.gz43_256998
M00054953B:D07	ES 199	555526	2005.J12.gz43_257062
M00054954A:F12	ES 199	561878	2005.K03.gz43_256919
M00054959A:A08	ES 199	556355	2005.O05.gz43_256955
M00054960C:C03	ES 199	456795	2006.A03.gz43_257293
M00054961A:E10	ES 199	550149	2006.A11.gz43_257421
M00054961B:D07	ES 199	556740	2006.A17.gz43_257517
M00054961B:G06	ES 199	562990	2006.A22.gz43_257597
M00054961C:H07	ES 199	450524	2006.B02.gz43_257278
M00054961D:A10	ES 199	556273	2006.B03.gz43_257294
M00054961D:H08	ES 199	557421	2006.B09.gz43_257390
M00054962A:H07	ES 199	129715	2006.B22.gz43_257598
M00054962B:A07	ES 199	555135	2006.B23.gz43_257614
M00054962C:D02	ES 199	599012	2006.C06.gz43_257343
M00054963A:C12	ES 199	531145	2006.C17.gz43_257519
M00054963A:E02	ES 199	560696	2006.C20.gz43_257567
M00054963A:G12	ES 199	553976	2006.C22.gz43_257599
M00054963A:H04	ES 199	471257	2006.C23.gz43_257615
M00054963B:H01	ES 199	449537	2006.D05.gz43_257328
M00054963C:C08	ES 199	555394	2006.D06.gz43_257344
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M00054964A:F09	ES 199	556954	2006.E03.gz43_257297
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M00054965C:C04	ES 199	557883	2006.F18.gz43_257538
M00054966A:C02	ES 199	465339	2006.G01.gz43 257267

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Table 13			
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M00054966A:D06	ES 199	593166	2006.G05.gz43_257331
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M00054966A:G02	ES 199	556126	2006.G10.gz43_257411
M00054966D:C05	ES 199	497938	2006.G22.gz43_257603
M00054966D:H12	ES 199	494306	2006.H03.gz43 257300
M00054967A:D11	ES 199	456203	2006.H05.gz43 257332
M00054967D:F08	ES 199	208278	2006.I03.gz43 257301
M00054968A:C11	ES 199	592941	2006.I06.gz43_257349
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M00054969A:E05	ES 199	556123	2006.J08.gz43 257382
M00054969A:F12	ES 199	554646	2006.J11.gz43 257430
M00054969B:F12	ES 199	556552	2006.J19.gz43_257558
M00054969C:G09	ES 199	556951	2006.J22.gz43 257606
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M00054973B:A10	ES 199	555103	2006.O06.gz43_257355
M00054973D:B09	ES 199	555155	2006.O16.gz43_257515
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M00054974D:C11	ES 199	556552	2006.P11.gz43_257436
M00054975A:C08	ES 199	559047	2006.P12.gz43_257452
M00054975A:E02	ES 199	552941	2006.P13.gz43_257468
M00054975B:B06	ES 199	554365	2006.P15.gz43_257500
M00054975C:D08	ES 199	549579	2006.P22.gz43_257612
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M00054976A:A03	ES 199	551967	2007.A05.gz43_257709
M00054976B:C10	ES 199	556512	2007.A08.gz43_257757
M00054976C:A03	ES 199	558670	2007.A11.gz43_257805
M00054976C:G10	ES 199	476517	2007.A19.gz43_257933
M00054976D:F06	ES 199	523332	2007.A24.gz43_258013
M00054977C:A09	ES 199	458257	2007.B13.gz43 257838

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M00054977C:F09	ES 199	561112	2007.B18.gz43_257918
M00054977D:B06	ES 199	551694	2007.B20.gz43_257950
M00054978C:E07	ES 199	556768	2007.C04.gz43_257695
M00054978D:H09	ES 199	562750	2007.C09.gz43_257775
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M00054979B:G12	ES 199	557054	2007.C18.gz43_257919
M00054980C:C07	ES 199	556603	2007.D09.gz43_257776
M00054980C:H08	ES 199	550333	2007.D11.gz43_257808
M00054980D:H07	ES 199	555197	2007.D13.gz43_257840
M00054981B:B09	ES 199	483410	2007.D18.gz43_257920
M00054981B:H12	ES 199	467293	2007.D21.gz43_257968
M00054981C:A11	ES 199	481243	2007.D24.gz43_258016
M00054981D:C03	ES 199	549822	2007.E06.gz43_257729
M00054981D:C06	ES 199	556561	2007.E08.gz43_257761
M00054982C:A02	ES 199	552437	2007.E15.gz43_257873
M00054983A:F08	ES 199	562655	2007.E20.gz43_257953
M00054983C:F05	ES 199	170450	2007.F04.gz43_257698
M00054983C:G09	ES 199	554123	2007.F05.gz43_257714
M00054983D:D05	ES 199	560652	2007.F09.gz43_257778
M00054984A:A05	ES 199	555512	2007.F10.gz43_257794
M00054984A:D12	ES 199	549581	2007.F12.gz43_257826
M00054985A:D09	ES 199	555594	2007.G10.gz43_257795
M00054985C:B12	ES 199	543429	2007.G17.gz43_257907
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M00054985D:E03	ES 199	560748	2007.G23.gz43_258003
M00054986B:D07	ES 199	555639	2007.H07.gz43_257748
M00054986C:D08	ES 199	553787	2007.H12.gz43_257828
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M00054987B:G05	ES 199	550814	2007.I05.gz43_257717
M00054987C:A11	ES 199	454463	2007.I08.gz43_257765
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M00054987C:G04	ES 199	556598	2007.I10.gz43_257797
M00054987D:D01	ES 199	475624	2007.I12.gz43_257829
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M00054988C:G02	ES 199	557039	2007.I19.gz43_257941
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M00054990C:A08	ES 199	556288	2007.J17.gz43 257910
M00054990C:E06	ES 199	554818	2007.J18.gz43_257926
M00054990D:A06	ES 199	556286	2007.J20.gz43_257958
M00054990D:F04	ES 200	503452	2007.J21.gz43_257974
M00054991B:E11	ES 200	556841	2007.K03.gz43_257687
M00054991C:E01	ES 200	454463	2007.K08.gz43_257767
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M00054992A:C08	ES 200	555277	2007.K19.gz43 257943
M00054992A:D11	ES 200	551798	2007.K20.gz43_257959
M00054992D:C12	ES 200	597957	2007.L03.gz43_257688
M00054992D:F06	ES 200	557811	2007.L05.gz43_257720
M00054993A:E04	ES 200	554885	2007.L07.gz43_257752
M00054993B:H06	ES 200	555658	2007.L15.gz43_257880
M00054993C:C10	ES 200	552055	2007.L17.gz43 257912
M00054993C:D12	ES 200	559372	2007.L20.gz43 257960
M00054993C:G12	ES 200	556019	2007.L22.gz43_257992
M00054993C:H05	ES 200	562323	2007.L23.gz43_258008
M00054993D:F04	ES 200	557420	2007.M02.gz43_257673
M00054994A:E05	ES 200	550833	2007.M06.gz43_257737
M00054994B:D11	ES 200	452682	2007.M11.gz43_257817
M00054994C:A01	ES 200	476398	2007.M13.gz43_257849
M00054994C:B12	ES 200	553252	2007.M14.gz43_257865
M00054994C:G06	ES 200	553797	2007.M16.gz43_257897
M00054995B:G12	ES 200	554764	2007.N06.gz43_257738
M00054995D:D10	ES 200	551150	2007.N13.gz43_257850
M00054996A:B01	ES 200	559514	2007.N17.gz43_257914
M00054996A:D10	ES 200	549214	2007.N20.gz43_257962
M00054996B:C11	ES 200	555958	2007.N24.gz43_258026
M00054997A:C02	ES 200	554212	2007.O12.gz43_257835
M00054997A:G11	ES 200	476732	2007.O15.gz43_257883
M00054997B:F10	ES 200	559096	2007.O18.gz43_257931
M00054997C:H03	ES 200	560984	2007.O24.gz43_258027
M00054997D:E01	ES 200	554395	2007.P06.gz43_257740
M00055000A:H10	ES 200	553915	2007.P11.gz43_257820
M00055000B:B04	ES 200	555883	2007.P12.gz43_257836
M00055000B:F02	ES 200	553732	2007.P13.gz43_257852
M00055000C:F08	ES 200	549516	2007.P17.gz43_257916
M00055000D:C03	ES 200	557531	2007.P19.gz43_257948
M00055001A:A02	ES 200	591128	2007.P24.gz43_258028
M00055001A:B10	ES 200	553064	2008.A02.gz43_258047
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Table 15			
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M00055001C:A11	ES 200	553967	2008.A15.gz43 258255
M00055001C:C10	ES 200	335714	2008.A17.gz43 258287
M00055001C:G07	ES 200	549690	2008.A18.gz43 258303
M00055002B:E07	ES 200	549912	2008.B03.gz43_258064
M00055002B:G03	ES 200	556042	2008.B05.gz43_258096
M00055002D:A07	ES 200	554379	2008.B11.gz43 258192
M00055002D:E04	ES 200	562550	2008.B12.gz43_258208
M00055003A:D05	ES 200	553191	2008.B15.gz43_258256
M00055003A:D11	ES 200	437580	2008.B16.gz43_258272
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M00055003A:H01	ES 200	601219	2008.B19.gz43_258320
M00055003B:E07	ES 200	555639	2008.B21.gz43_258352
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M00055003D:F09	ES 200	134237	2008.C09.gz43_258161
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M00055005D:C11	ES 200	555289	2008.D18.gz43_258306
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M00055006B:A09	ES 200	558212	2008.E05.gz43_258099
M00055006B:C08	ES 200	552674	2008.E07.gz43_258131
M00055006B:E05	ES 200	347057	2008.E10.gz43_258179
M00055006B:G03	ES 200	557026	2008.E11.gz43_258195
M00055007C:D01	ES 200	34381	2008.F08.gz43_258148
M00055007D:D11	ES 200	480960	2008.F13.gz43_258228
M00055008B:E08	ES 200	550497	2008.F18.gz43_258308
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M00055010B:A06	ES 200	460169	2008.H22.gz43_258374
M00055010B;C11	ES 200	497513	2008.H24.gz43_258406
M00055010C:C10	ES 200	597780	2008.I06.gz43_258119
M00055010D:A02	ES 200	555371	2008.I09.gz43_258167
M00055010D:D08	ES 200	555512	2008.I13.gz43_258231
M00055010D:E08	ES 200	555658	2008.I14.gz43_258247
M00055010D:F11	ES 200	600021	2008.I15.gz43_258263
M00055011A:B11	ES 200	362109	2008.I17.gz43_258295
M00055011A:C06	ES 200	559464	2008.I18.gz43 258311
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ES 200	551268	2008.O16.gz43_258283
ES 200	556488	2008.P01.gz43_258044
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ES 200	556115	2008.P04.gz43_258094
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ES 200	556487	2008.P11.gz43_25820
	556490	2008.P12.gz43_258222
		2008.P20.gz43_258350
	608873	2017.A02.gz43_25843
	557910	2017.B02.gz43_25843
		2017.B05.gz43_25848
		2017.B14.gz43_25862
		2017.B22.gz43_25875
		2017.C09.gz43_25854
		2017.C14.gz43_25862
		2017.D08.gz43_25853.
		2017.D10.gz43_25856
		2017.D11.gz43_25858
		2017.D14.gz43_25862
		2017.D18.gz43_25869
		2017 D19 gz43_25870
		2017,D21.gz43_25874
		2017.D22.gz43_25875
		2017.E05.gz43_25848
		2017.E06.gz43_25850
		2017.E07.gz43_25851
		2017.E10.gz43_25856
		2017.E11.gz43_25858
		2017.F06.gz43_25850
		2017.F16.gz43_25866
		2017.F19.gz43_25871
		2017.F21.gz43_25874
		2017,F22,gz43_25875
		2017.G01.gz43_25842
		2017.G09.gz43_25855
		2017.G16.gz43_25866
		2017.H04.gz43_258472 2017.H06.gz43_25850
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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055028D:E10	ES 200	560294	2017.H10.gz43_258568
M00055029A:A02	ES 200	560621	2017.H11.gz43_258584
M00055029C:D05	ES 200	557603	2017.I02.gz43_258441
M00055030B;E03	ES 200	549599	2017.I11.gz43_258585
M00055030B;H07	ES 200	461850	2017.I14.gz43_258633
M00055031B:E02	ES 200	561301	2017.J06.gz43_258506
M00055031B:E05	ES 200	557733	2017.J08.gz43_258538
M00055031B:H08	ES 200	- 558004	2017.J11.gz43_258586
M00055031C:B11	ES 200	552848	2017.J14.gz43_258634
M00055032B:D08	ES 200	561279	2017.K06.gz43 258507
M00055032D:B06	ES 200	454014	2017,K16.gz43 258667
M00055033B;H01	ES 200	562590	2017.L12.gz43 258604
M00055033C:A08	ES 200	557283	2017.L13.gz43 258620
M00055033D:D03	ES 200	557010	2017.L19.gz43 258716
M00055033D:G01	ES 200	557883	2017.L21.gz43 258748
M00055034A:G01	ES 200	460078	2017.L23.gz43_258780
M00055034B:G04	ES 200	523495	2017.M03.gz43_258461
M00055034D:A10	ES 200	556458	2017,M13.gz43_258621
M00055035B:D11	ES 200	408586	2017.M22.gz43_258765
M00055035C:G10	ES 200	555651	2017.N07.gz43_258526
M00055035D:E07	ES 200	556852	2017.N09.gz43_258558
M00055035D:F09	ES 200	558755	2017.N12.gz43_258606
M00055036A:H10	ES 200	456183	2017.N18.gz43_258702
M00055036B:H02	ES 201	561836	2017.N23.gz43_258782
M00055036C:G10	ES 201	491127	2017.O02.gz43_258447
M00055037C:D01	ES 201	557578	2017.O18.gz43_258703
M00055038A:H08	ES 201	134734	2017.P07.gz43_258528
M00055039A:G06	ES 201	561438	2018.A03.gz43_264190
M00055039B:E02	ES 201	555660	· 2018.A07.gz43_264254
M00055039B:G11	ES 201	555993	2018.A08.gz43_264270
M00055039B:G11	ES 201	555993	2018.A08.gz43_264654
M00055039B:H10	ES 201	601365	2018.A09.gz43_264286
M00055039C:B05	ES 201	529733	2018.A11.gz43 264318
M00055039C:D11	ES 201	397338	2018.A12.gz43_264718
M00055039D:D07	ES 201	. 561259	2018.A15.gz43_264382
M00055039D:D07	ES 201	561259	2018.A15.gz43 264766
M00055040A:C02	ES 201	555399	2018,A17.gz43_264414
M00055040A:C02	ES 201	555399	2018.A17.gz43_264798
M00055040A:F01	ES 201	555883	2018.A18.gz43_264430
M00055040B:F02	ES 201	555751	2018.A22.gz43_264878
M00055040C:G08	ES 201	447815	2018.A23.gz43_264510
M00055040D:B05	ES 201	555160	2018.A24.gz43_264910
M00055040D:G05	ES 201	561761	2018.B02.gz43_264175

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055041A:C02	ES 201	555340	2018.B05.gz43_264223
M00055041B:B06	ES 201	417274	2018.B08.gz43_264655
M00055041B:C10	ES 201	551711	2018.B09.gz43_264671
M00055041B:D11	ES 201	555524	2018,B10,gz43_264303
M00055041B:D11	ES 201	555524	2018.B10.gz43_264687
M00055041B:F04	ES 201	561994	2018.B13.gz43_264735
M00055041C:C10	ES 201	554149	2018.B17.gz43_264799
M00055041C:H12	ES 201	470617	2018.B22.gz43_264495
M00055041D:B07	ES 201	555250	2018.B23.gz43_264895
M00055042A:D09	ES 201	561901	2018.C07.gz43_264256
M00055042A:F07	ES 201	551328	2018.C08.gz43_264656
M00055042B:B09	ES 201	549008	2018.C10.gz43_264304
M00055042B:E02	ES 201	555708	2018.C12.gz43_264336
M00055042B:E05	ES 201	448098	2018.C13.gz43_264736
M00055042B:E08	ES 201	450765	2018.C14.gz43_264368
M00055042B:E08	ES 201	450765	2018.C14.gz43_264752
M00055042C:B05	ES 201	555172	2018.C16.gz43 264400
M00055042C:F04	ES 201	551805	2018.C19.gz43 264832
M00055042D:G03	ES 201	555940	2018.C21.gz43 264864
M00055042D:H02	ES 201	423588	2018.C22.gz43_264880
M00055043A:B06	ES 201	555173	2018.C24.gz43 264912
M00055043B:B01	ES 201	555213	2018,D07.gz43 264641
M00055043B:G01	ES 201	551714	2018.D08.gz43 264273
M00055043D:D10	ES 201	558254	2018.D13.gz43 264737
M00055043D:F07	ES 201	553356	2018.D14.gz43 264753
M00055044A:A08	ES 201	562949	2018.D17.gz43_264417
M00055044A:C02	ES 201	450755	2018.D18.gz43_264817
M00055044B:F12	ES 201	549444	2018.D21.gz43_264865
M00055045A:C06	ES 201	561911	2018.E08.gz43_264658
M00055045A:F03	ES 201	555771	2018.E10.gz43_264306
M00055045A:F12	ES 201	553850	2018.E11.gz43_264322
M00055045B:A04	ES 201	559699	2018.E12.gz43_264338
M00055045B:A04	ES 201	559699	2018.E12.gz43_264722
M00055045B:A12	ES 201	347486	2018.E13.gz43_264738
M00055045B:C08	ES 201	560720	2018.E14.gz43_264370
M00055045B:C08	ES 201	560720	2018.E14.gz43_264754
M00055045C:F09	ES 201	555892	2018.E18.gz43_264818
M00055045C:H05	ES 201	556158	2018.E19.gz43 264450
M00055045D;A01	ES 201	108755	2018.E21.gz43_264482
M00055045D:A07	ES 201	551578	2018.E22.gz43_264882
M00055045D:D01	ES 201	446664	2018.E24.gz43_264914
M00055045D:F10	ES 201	561838	2018.F02.gz43_264179
M00055046A:D06	ES 201	562585	2018.F05.gz43_264227
M00055046B:C07	ES 201	375380	2018.F09.gz43_264291

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Table 15			
CloneID	ES No	ClusterID	SequenceName
M00055046B:C12	ES 201	557546	2018.F10.gz43_264307
M00055046B:D02	ES 201	559052	2018.F11.gz43_264707
M00055046B:E08	ES 201	557420	2018.F13.gz43_264739
M00055046B:F06	ES 201	491799	2018.F14.gz43_264755
M00055046C:C05	ES 201	558389	2018.F18.gz43_264435
M00055046C:E07	ES 201	488970	2018.F21.gz43_264867
M00055046C:E11	ES 201	514418	2018.F23.gz43_264515
M00055046C:G10	ES 201	551641	2018.F24.gz43_264915
M00055047A:E07	ES 201	557735	2018.G11.gz43_264324
M00055047A:H04	ES 201	558007	2018.G13.gz43_264740
M00055047A:H05	ES 201	558120	2018.G14.gz43_264372
M00055047A:H05	ES 201	558120	2018.G14.gz43_264756
M00055047B:A10	ES 201	557326	2018.G15.gz43_264388
M00055047B:B03	ES 201	289316	2018.G16.gz43_264404
M00055047B:B10	ES 201	553603	2018.G17.gz43_264804
M00055047B:C03	ES 201	551793	2018.G18.gz43_264820
M00055047B:G06	ES 201	556171	2018.G22.gz43_264884
M00055047B:G10	ES 201	476268	2018.G23.gz43 264900
M00055047C:D11	ES 201	447455	2018.H06.gz43 264245
M00055047C:F07	ES 201	554936	2018.H07.gz43_264261
M00055047C:F08	ES 201	465207	2018.H08.gz43_264277
M00055047C:F08	ES 201	465207	2018.H08.gz43_264661
M00055047D:C12	ES 201	555798	2018.H11.gz43_264325
M00055048A;A04	ES 201	557279	2018.H16.gz43_264405
M00055048A:B12	ES 201	465447	2018.H17.gz43_264421
M00055048A:D12	ES 201	557606	2018.H19.gz43_264453
M00055048A:F04	ES 201	453893	2018.H20.gz43_264469
M00055048C:C06	ES 201	556490	2018.I04.gz43_264214
M00055048D:D08	ES 201	419153	2018.I08.gz43_264278
M00055048D:D08	ES 201	419153	2018.I08.gz43_264662
M00055048D:H04	ES 201	349744	2018.I14.gz43_264374
M00055049A:F10	ES 201	559762	2018.I19.gz43_264838
M00055049A:G03	ES 201	561046	2018.I20.gz43_264854
M00055049B:A01	ES 201	557249	2018.I22.gz43_264502
M00055049B:F05	ES 201	553983	2018.J01.gz43_264167
M00055049C:H12	ES 201	555010	2018.J07.gz43_264263
M00055049C:H12	ES 201	555010	2018.J07.gz43_264647
M00055049D:D09	ES 201	561216	2018.J09.gz43_264295
M00055049D:D10	ES 201	446739	2018.J10.gz43_264695
M00055050A:D11	ES 201	449437	2018.J15.gz43_264391
M00055050A:H08	ES 201	556169	2018.J16.gz43_264407
M00055050B;E11	ES 201	206098	2018.J19.gz43_264455
M00055050B;E11	ES 201	206098	2018.J19.gz43_264839
M00055050D:C01	ES 201	559775	2018.K06.gz43_264248

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CloneID	ES No	ClusterID	SequenceName
M00055050D:E03	ES 201	486150	2018.K07.gz43_264264
M00055050D:F04	ES 201	557834	2018.K08.gz43_264664
M00055051A:B02	ES 201	454810	2018.K11.gz43_264328
M00055051A:C09	ES 201	551068	2018.K12.gz43_264344
M00055051A:E11	ES 201	432970	2018.K14.gz43_264760
M00055051A:G09	ES 201	555340	2018.K15.gz43_264392
M00055051A:G09	ES 201	555340	2018.K15.gz43_264776
M00055051A:H10	ES 201	551897	2018.K16.gz43_264408
M00055051B;B08	ES 201	552641	2018.K17.gz43_264424
M00055051B:B08	ES 201	552641	2018.K17.gz43_264808
M00055051B:D07	ES 201	557572	2018.K19.gz43_264840
M00055051B:G09	ES 201	451027	2018.K21.gz43_264872
M00055051C:B05	ES 201	553349	2018.K23.gz43_264904
M00055051C:F10	ES 201	554372	2018.L04.gz43_264217
M00055051D:D12	ES 201	540000	2018.L05.gz43_264233
M00055051D:F01	ES 201	560801	2018.L07.gz43_264265
M00055051D:G01	ES 201	557882	2018.L08.gz43_264665
M00055052A:F07	ES 201	559246	2018.L12.gz43_264345
M00055052A:H11	ES 201	558029	2018.L14.gz43_264761
M00055052B:E03	ES 201	551371	2018.L17.gz43_264425
M00055052B;E03	ES 201	551371	2018.L17.gz43_264809
M00055052C:B12	ES 201	454664	2018.L20.gz43_264857
M00055052D:B05	ES 201	557382	2018.L24.gz43_264921
M00055052D:G12	ES 201	493261	2018.M06.gz43_264250
M00055053B:A02	ES 201	553108	2018.M11.gz43_264330
M00055053B:A02	ES 201	553108	2018.M11.gz43_264714
M00055053B;C02	ES 201	557525	2018.M13.gz43_264362
M00055053B:C02	ES 201	557525	2018.M13.gz43_264746
M00055053B;C11	ES 201	558560	2018.M14.gz43_264762
M00055053C:A12	ES 201	557345	2018.M16.gz43_264794
M00055053C:B03	ES 201	562292	2018,M17.gz43_264426
M00055053C:B03	ES 201	562292	2018,M17.gz43_264810
M00055053C:F06	ES 201	551334	2018.M19.gz43 264842
M00055053D:G04	ES 201	557939	2018.N01.gz43_264171
M00055054A:A10	ES 201	447555	2018.N03.gz43 264203
M00055054B:E10	ES 201	8997	2018.N13.gz43 264363
M00055054B:E10	ES 201	8997	2018.N13.gz43 264747
M00055054B:F05	ES 201	562115	2018.N15.gz43 264395
M00055054B:G12	ES 201	555082	2018.N17.gz43 264427
M00055054C:G10	ES 201	551616	2018.N20.gz43_264859
M00055054D:A02	ES 201	557308	2018.N21.gz43_264491
M00055054D:E12	ES 201	555057	2018.N22.gz43 264507
M00055054D:H07	ES 201	467081	2018.N23.gz43 264523
M00055054D:H07	ES 201	467081	2018.N23.gz43 264907

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M00055055A:B03	
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M00055056C:F04 ES 201 M00055056C:H07 ES 201 M00055056C:H07 ES 201 M00055056D:B06 ES 201 M00055056D:H12 ES 201 M00055056D:H12 ES 201 M00055057A:A04 ES 201 M00055057A:A05 ES 201 M00055057A:A05 ES 201 M00055057A:A05 ES 201 M00055057A:B05 ES 201 M00055057A:B10 ES 201 M00055057A:B10 ES 201 M00055057A:B10 ES 201 M00055057A:B10 ES 201 M00055057A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201 M0005507A:B10 ES 201	553012 2018.P05.gz43_26423
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M00055056D:B06 ES 201 M00055056D:H12 ES 201 M00055056D:H12 ES 201 M00055057A:A04 ES 201 M00055057A:A05 ES 201 M00055057A:A05 ES 201 M00055057A:D10 ES 201 M00055057A:D10 ES 201 M00055057A:D10 ES 201 M00055057A:D10 ES 201 M00055057A:D10 ES 201 M0005507AB:B01 ES 201 M0005507AB:B01 ES 201 M0005507AB:B01 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B05 ES 201 M0005507AB:B08 ES 201 M0005507AB:B08 ES 201 M0005507AB:B08 ES 201 M0005507AB:B08 ES 201 M0005507AB:B08 ES 201 M0005507AB:B08 ES 201 M0005507AB:B08 ES 201	562808 2018.P08.gz43_26428
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M00055057A:A05 ES 201 M00055057A:D10 ES 201 M00055057A:D10 ES 201 M00055057A:B10 ES 201 M00055057B:B01 ES 201 M00055073B:B01 ES 201 M00055074B:E05 ES 201 M00055074B:E05 ES 201 M00055074B:B04 ES 201 M00055075A:C09 ES 201 M00055075A:C09 ES 201 M00055075A:C00 ES 201 M00055076A:C06 ES 201 M00055076A:C06 ES 201 M00055076A:C06 ES 201 M00055076A:D11 ES 201 M00055076A:G12 ES 201 M00055076A:G12 ES 201 M00055076B:E08 ES 201 M00055076B:E08 ES 201 M00055076B:E08 ES 201 M00055076B:E08 ES 201 M00055076B:E08 ES 201 M00055076B:E08 ES 201 M00055076B:E08 ES 201 M00055076B:E08 ES 201 M00055076B:E08 ES 201 M00055076B:E08 ES 201 M00055076B:E08 ES 201 M00055076B:E08 ES 201	562272 2018.P17.gz43_26481
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M00055073D:F06 ES 201 M00055074B:E05 ES 201 M00055074B:E05 ES 201 M00055074B:D05 ES 201 M00055075A:C09 ES 201 M00055075B:D12 ES 201 M00055075CA:C06 ES 201 M00055076A:C06 ES 201 M00055076A:C01 ES 201 M00055076A:C01 ES 201 M00055076A:F04 ES 201 M00055076B:E08 ES 201 M00055076B:F04 ES 201 M00055076B:F06 ES 201 M00055076B:F06 ES 201 M00055076B:F06 ES 201 M00055076B:F06 ES 201 M00055076B:F06 ES 201 M00055076B:F06 ES 201	451172 2018.P22.gz43_26489
M00055074B:E05 ES 201 M00055074D:B04 ES 201 M00055075A:C09 ES 201 M00055075B:D12 ES 201 M00055075B:D12 ES 201 M00055075D:D05 ES 201 M00055076A:C06 ES 201 M00055076A:D11 ES 201 M00055076A:D11 ES 201 M00055076B:E08 ES 201 M00055076B:F04 ES 201 M00055076B:F04 ES 201 M00055076B:F06 ES 201 M00055076B:F06 ES 201	551544 2018.P24.gz43_26454
M00055074D:B04 ES 201 M00055075A:C09 ES 201 M00055075B:D12 ES 201 M00055075D:D05 ES 201 M00055076A:C06 ES 201 M00055076A:C11 ES 201 M00055076A:G12 ES 201 M00055076A:G12 ES 201 M00055076B:E08 ES 201 M00055076B:E08 ES 201 M00055076B:F04 ES 201 M00055076B:F04 ES 201 M00055076B:F06 ES 201 M00055076B:F06 ES 201 M00055076B:F06 ES 201	555512 2020.A04.gz43_26497
M00055075A:C09 ES 201 M00055073B:D12 ES 201 M00055075D:D05 ES 201 M00055076A:C06 ES 201 M00055076A:C10 ES 201 M00055076A:G12 ES 201 M00055076B:E08 ES 201 M00055076B:F04 ES 201 M00055076B:F06 ES 201 M00055076B:F06 ES 201 M00055076B:F06 ES 201 M00055076B:F06 ES 201 M00055076B:F06 ES 201	455820 2020.A14.gz43_26513
M00055075B:D12 ES 201 M00055075D:D05 ES 201 M00055076A:C06 ES 201 M00055076A:C06 ES 201 M00055076A:D11 ES 201 M00055076B:E08 ES 201 M00055076B:E08 ES 201 M00055076B:F04 ES 201 M00055076B:F04 ES 201 M00055076B:F06 ES 201	562229 2020,A22,gz43_26526
M00055075D:D05 ES 201 M00055076A:C06 ES 201 M00055076A:G11 ES 201 M00055076A:G12 ES 201 M00055076B:E08 ES 201 M00055076B:F04 ES 201 M00055076B:F04 ES 201 M00055076B:H06 ES 201 M00055076B:H06 ES 201	556635 2020.B05.gz43_26499
M00055076A:C06 ES 201 M00055076A:D11 ES 201 M00055076A:G12 ES 201 M00055076B:E08 ES 201 M00055076B:F04 ES 201 M00055076B:F06 ES 201 M00055076B:H06 ES 201 M00055076C:B10 ES 201	556742 2020.B12.gz43_26510
M00055076A:D11 ES 201 M00055076A:G12 ES 201 M00055076B:E08 ES 201 M00055076B:F04 ES 201 M00055076B:F04 ES 201 M00055076B:H06 ES 201 M00055076C:B10 ES 201	560069 2020.B20.gz43_26523
M00055076A:G12 ES 201 M00055076B:E08 ES 201 M00055076B:F04 ES 201 M00055076B:H06 ES 201 M00055076C:B10 ES 201	558720 2020.C03.gz43_26496
M00055076B:E08 ES 201 M00055076B:F04 ES 201 M00055076B:H06 ES 201 M00055076C:B10 ES 201	555368 2020,C05.gz43_26499
M00055076B:F04 ES 201 M00055076B:H06 ES 201 M00055076C:B10 ES 201	557024 2020.C07.gz43_26502
M00055076B:H06 ES 201 M00055076C:B10 ES 201	452822 2020.C11.gz43_26508
M00055076C:B10 ES 201	549810 2020.C12.gz43_26510
	550701 2020.C16.gz43_26516
	559965 2020.C18.gz43_26520
	557954 2020,C20.gz43_26523
	557783 2020, C23, gz43_26528
M00055076D:H11 ES 201	553131 2020,D02,gz43_26494

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Table 13

Table 15			
CloneID	ES No	ClusterID	SequenceName
M00055077A:B07	ES 201	549129	2020.D04.gz43_264977
M00055077A:H10	ES 201	561707	2020.D10.gz43 265073
M00055077D:B01	ES 201	556471	2020.E02.gz43_264946
M00055078A:C05	ES 201	142614	2020.E08.gz43 265042
M00055078A:E10	ES 201	464905	2020,E11.gz43_265090
M00055078A:F01	ES 201	557760	2020.E12.gz43_265106
M00055078A:F04	ES 201	553303	2020.E13.gz43_265122
M00055078B:F05	ES 201	561489	2020.E15.gz43_265154
M00055078D:A07	ES 201	555996	2020.E21.gz43_265250
M00055078D:G04	ES 201	495408	2020.F06.gz43_265011
M00055079A:H05	ES 201	509505	2020.F11.gz43_265091
M00055079B:D02	ES 201	402683	2020.F12.gz43_265107
M00055079C:G06	ES 201	558616	2020.F15.gz43_265155
M00055079D:A03	ES 201	612961	2020.F17.gz43_265187
M00055080A:A07	ES 201	555111	2020.F22.gz43_265267
M00055080A:F05	ES 201	451185	2020.F24.gz43_265299
M00055080B:G10	ES 201	556040	2020.G09.gz43_265060
M00055080D:A01	ES 201	551976	2020.G16.gz43_265172
M00055080D:E07	ES 201	553318	2020,G19,gz43_265220
M00055080D:E10	ES 201	556357	2020.G20.gz43_265236
M00055080D:F01	ES 201	494625	2020.G21.gz43_265252
M00055081A:E08	ES 202	555616	2020.H01.gz43_264933
M00055081B:E10	ES 202	552361	2020.H06.gz43_265013
M00055081C:A12	ES 202	555061	2020.H11.gz43_265093
M00055081C:G01	ES 202	553372	2020.H18.gz43_265205
M00055081C:H04	ES 202	449613	2020.H19.gz43_265221
M00055082A:A12	ES 202	552019	2020.H24.gz43_265301
M00055082D:E08	ES 202	555710	2020.I14.gz43_265142
M00055082D:G01	ES 202	460244	2020,I16.gz43_265174
M00055082D:H02	ES 202	465446	2020.I17.gz43_265190
M00055083B:E05	ES 202	549607	2020.J01.gz43_264935
M00055083B:E07	ES 202	452434	2020.J02.gz43_264951
M00055083B;F10	ES 202	552618	2020.J03.gz43_264967
M00055083C:C05	ES 202	556497	2020.J04.gz43_264983
M00055083C:F05	ES 202	556882	2020.J05.gz43_264999
M00055083D:D08	ES 202	556668	2020.J11.gz43_265095
M00055084A;E10	ES 202	556802	2020.J16.gz43_265175
M00055084A;F10	ES 202	556925	2020.J17.gz43_265191
M00055084B:A04	ES 202	558231	2020.J18.gz43_265207
M00055084D:B01	ES 202	556446	2020.K06.gz43_265016
M00055084D:C09	ES 202	548943	2020,K07.gz43_265032
M00055085A:F12	ES 202	500833	2020.K12.gz43_265112
M00055085B:D02	ES 202	454910	2020.K16.gz43_265176

Table 13

CloneID	ES No	ClusterID	SequenceName
M00055085C:E05	ES 202	561753	2020.K19.gz43_265224
M00055085D:D10	ES 202	451134	2020.K21.gz43 265256
M00055085D:F03	ES 202	556881	2020:K23.gz43_265288
M00055085D:G09	ES 202	555725	2020.K24.gz43 265304
M00055086A:B10	ES 202	492893	2020.L01.gz43 264937
M00055086A:C09	ES 202	556542	2020,L02.gz43_264953
M00055086B:D10	ES 202	557476	2020.L06.gz43 265017
M00055086D:C07	ES 202	551342	2020.L13.gz43 265129
M00055086D:H07	ES 202	557214	2020.L16.gz43_265177
M00055087B:C03	ES 202	122169	2020.L22.gz43_265273
M00055087B:C05	ES 202	451885	2020.L23.gz43_265289
M00055087D:D08	ES 202	453756	2020,M08.gz43_265050
M00055088C:B02	ES 202	450352	2020.M20.gz43_265242
M00055088C:D01	ES 202	561975	2020.M22.gz43 265274
M00055088C:D02	ES 202	557747	2020.M23.gz43_265290
M00055088D:A01	ES 202	551283	2020.N02.gz43_264955
M00055088D:B08	ES 202	561422	2020,N04.gz43 264987
M00055089A:H06	ES 202	557454	2020.N09.gz43_265067
M00055089B:C01	ES 202	554084	2020.N12.gz43_265115
M00055089C:B06	ES 202	556465	2020.N16.gz43_265179
M00055089C:D06	ES 202	560507	2020.N19.gz43_265227
M00055090A:F02	ES 202	560080	2020.N22.gz43_265275
M00055090C:B03	ES 202	556408	2020.O04.gz43_264988
M00055090C:C12	ES 202	556559	2020.O06.gz43_265020
M00055090C:D01	ES 202	556675	2020.O07.gz43_265036
M00055090C:G06	ES 202	451370	2020.O09.gz43_265068
M00055090D:E03	ES 202	497086	2020.O14.gz43_265148
M00055090D:F03	ES 202	98869	2020.O17.gz43_265196
M00055091A:C09	ES 202	598101	2020,O23.gz43_265292
M00055091A:F05	ES 202	459581	2020.P04.gz43_264989
M00055091B:A07	ES 202	555126	2020.P07.gz43_265037
M00055091B;C11	ES 202	555418	2020,P08.gz43_265053
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M00055091C:D11	ES 202	559071	2020.P15.gz43_265165
M00055091C:G11	ES 202	561593	2020,P18.gz43_265213
M00055091D:A03	ES 202	554828	2020.P21.gz43_265261
M00055092B:G09	ES 202	637966	2029.A15.gz43_265534
M00055093A;E09	ES 202	562542	2029.B05.gz43_265375
M00055093A:F07	ES 202	562768	2029.B06.gz43_265391
M00055093B:G08	ES 202	562881	2029.B14.gz43_265519
M00055094B:B11	ES 202	562307	2029.C11.gz43_265472
M00055094C:C10	ES 202	402488	2029.C21.gz43_265632
M00055094D:F09	ES 202	491127	2029.D06.gz43_265393
M00055095A;D08	ES 202	98484	2029.D11.gz43_265473

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Table 13

Table 13 CloneID	ES No	ClusterID	SequenceName
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M00055096D:F02	ES 202	553380	2029,E22.gz43 265650
M00055097A:G06	ES 202	455820	2029,F06.gz43 265395
M00055097B:B12	ES 202	453533	2029.F09.gz43 265443
M00055097B:F08	ES 202	562719	2029.F10.gz43 265459
M00055098D:A09	ES 202	561215	2029.G03.gz43 265348
M00055099A;G05	ES 202	558118	2029.G11.gz43 265476
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M00055100B:F11	ES 202	558186	2029.H15.gz43 265541
M00055100C:E03	ES 202	559125	2029.H22.gz43 265653
M00055100C:F11	ES 202	452349	2029.I01.gz43 265318
M00055100C:H06	ES 202	562989	2029.I02.gz43 265334
M00055100D:B02	ES 202	562243	2029.I03.gz43 265350
M00055102A:E11	ES 202	560628	2029.I20.gz43 265622
M00055102B:B03	ES 202	560111	2029.I21.gz43_265638
M00055103A:H11	ES 202	561144	2029.J16.gz43 265559
M00055103C:B07	ES 202	558511	2029.J20.gz43 265623
M00055103C:D05	ES 202	556613	2029.J21.gz43_265639
M00055103C:G03	ES 202	560898	2029.J22.gz43_265655
M00055104B:F09	ES 202	553548	2029.K14.gz43_265528
M00055104D:E02	ES 202	455878	2029.K20.gz43_265624
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M00055111C:B07	ES 202	558186	2029.P05.gz43_265389
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M00055113B:F02	ES 202	562714	2030,A12.gz43_265870
M00055113C:F09	ES 202	551734	2030,A14.gz43_265902
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M00055117C:F02	ES 202	561265	2030.D17.gz43_265953
M00055118B:A09	ES 202	554737	2030.D24.gz43_266065

Table 13

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M00055124D:H11	S 202	560506	2030.E22.gz43 266034
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M00055128B:B08 E: M00055128B:B12 E: M00055128B:B12 E: M00055128B:B12 E: M00055128B:G01 E: M00055128C:B03 E: M00055128C:B03 E: M00055128D:D04 E: M00055129B:B03 E: M00055129B:B03 E: M00055129B:B03 E: M00055129B:B03 E: M00055129B:B03 E: M00055129B:B03 E: M00055129B:B03 E: M00055129B:B03 E: M00055129B:B03 E: M00055129B:B03 E: M00055129B:B03 E: M00055129B:B03 E: M00055129B:B03 E: M00055129B:B03 E: M00055129B:B03 E: M00055129B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:B03 E: M00055130B:M00055130B:M00055130B:M00055130B:M00055130B:M00055130B:M00055130B:M00055130B:M00055130B:M00055130B:M00055130B:M00055130B:M00055130B:M00055130B:M00055130B:M00055130B:M00055130B:M0005	S 202	559938	2030.I12.gz43_265878
M00055128B:B12 EX M00055128B:B12 EX M00055128B:G1 EX M00055128C:E03 EX M00055128C:E03 EX M00055128C:E03 EX M00055129C:E03 EX M00055129C:E03 EX M00055129C:E03 EX M00055129C:E03 EX M00055129C:E03 EX M00055129C:E03 EX M00055129C:E03 EX M00055129C:E03 EX M00055129C:E03 EX M00055129C:E03 EX M00055129C:E03 EX M00055129C:E03 EX M00055139C:E03 EX M00055130A:E03 EX M00055130B:E0	S 202	554233	2030.J01.gz43_265703
M00055128B:E12 EX M00055128B:G01 EX M00055128B:G01 EX M00055128D:C11 EX M00055128D:D14 EX M00055129D:D14 EX M00055129D:H07 EX M00055129C:H08 EX M00055129C:H08 EX M00055129C:H08 EX M00055129C:H08 EX M00055129C:H08 EX M00055129C:H08 EX M00055129C:H08 EX M00055129C:H08 EX M00055129C:H08 EX M00055129D:F11 EX M00055130D:D10 EX M00055130B:D	S 202	561679	2030.J06.gz43_265783
M00055128B;G01 E8 M00055128C:E03 E8 M00055128D:D04 E8 M00055128D:D04 E8 M00055129B;B03 E8 M00055129B;B07 E8 M00055129C:H08 E8 M00055129C:H08 E8 M00055129C:H08 E8 M00055129D:F11 E8 M00055130A:D10 E8 M00055130B:D07 E8	S 202	528369	2030,J07.gz43_265799
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M00055128D:C11 ES M00055128D:D04 ES M00055129B:H07 ES M00055129C:H08 ES M00055129D:C02 ES M00055129D:C11 ES M00055129D:F11 ES M00055129D:F11 ES M00055130B:D07 ES	S 202	407964	2030.J13.gz43_265895
M00055128D:D04 ES M00055129A:B03 ES M00055129B:H07 ES M00055129C:H08 ES M00055129D:C02 ES M00055129D:F11 ES M00055130B:D07 ES	S 202	554989	2030.J16.gz43_265943
M00055129A:B03 ES M00055129B:H07 ES M00055129C:H08 ES M00055129D:C02 ES M00055129D:F11 ES M00055130A:D10 ES M00055130B:D07 ES	S 202	561279	2030,J22.gz43_266039
M00055129B:H07 ES M00055129C:H08 ES M00055129D:C02 ES M00055129D:F11 ES M00055130A:D10 ES M00055130B:D07 ES	S 202	557615	2030.K01.gz43_265704
M00055129C:H08 ES M00055129D:C02 ES M00055129D:F11 ES M00055130A:D10 ES M00055130B:D07 ES	S 202	486683	2030.K05.gz43_265768
M00055129D:C02 ES M00055129D:F11 ES M00055130A:D10 ES M00055130B:D07 ES	S 202	558024	2030.K18.gz43_265976
M00055129D:F11 ES M00055130A:D10 ES M00055130B:D07 ES	S 202 .	452759	2030.K24.gz43_266072
M00055130A:D10 ES M00055130B:D07 ES	S 202	452775	2030.L03.gz43_265737
M00055130B:D07 ES	S 202	488030	2030.L06.gz43_265785
	S 202	557610	2030.L11.gz43_265865
	S 202	380636	2030.L17.gz43_265961
M00055130D:F08 ES	S 202	66678	2030.M06.gz43_265786
M00055131A:A04 E	S 202	518007	2030.M10.gz43_265850
	S 202	449454	2030.M13.gz43_265898
M00055131C:H12 ES	S 202	562822	2030.N04.gz43_265755
M00055132B;B12 E	S 202	288134	2030.N12.gz43_265883
	S 202	610269	2030.N21.gz43_266027
	S 202	557719	2030.N22.gz43_266043
M00055132D:F05 ES	S 202	548991	2030.N23.gz43_266059

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Table 13

M00055133B:B06	Table 15			
M00055133B:E04		ES No	ClusterID	SequenceName
M00055133B:E08				2030.O05 gz43_265772
M00055133B:F07			557741	2030.O07.gz43_265804
M00055133C:C06		ES 202	557747	2030.O08.gz43_265820
M00055133C.G07	M00055133B:F07	ES 202	451618	2030.O10.gz43_265852
M00055133C:H11	M00055133C:C06	ES 202	562760	2030.O13.gz43_265900
M00055133D:A02	M00055133C:G07	ES 202	451392	2030.O15.gz43_265932
M00055133D:F02	M00055133C:H11	ES 202	558015	2030.O17.gz43_265964
M00055134A;C03	M00055133D:A02	ES 202	477295	2030.O18.gz43_265980
M00055134C:A01		ES 202	451391	2030,O22.gz43_266044
M00055134C:B01	M00055134A:C03	ES 202	553871	2030.O24.gz43_266076
M00055134C:E09	M00055134C:A01	ES 202	557284	2030.P11.gz43_265869
M00055134D:B03	M00055134C:B01	ES 202	551068	2030,P12.gz43_265885
M00055135A:D08	M00055134C:E09	ES 202	557710	2030.P13.gz43_265901
M00055138A:B07	M00055134D:B03	ES 202	407077	2030.P17.gz43_265965
M00055138A:H10	M00055135A:D08	ES 202	557549	2030.P23.gz43_266061
M00055136C:P11	M00055135A:E07	ES 202	562683	2031.A01.gz43_266078
M00055136D:D09	M00055135A:H10	ES 202	562725	2031.A02.gz43_266094
M00055137A:B05	M00055136C:F11	ES 202	562449	2031.A24.gz43_266446
M00055137B:B11	M00055136D:D09	ES 202	558768	2031,B05,gz43_266143
M00055137B:F12	M00055137A:E05	ES 202	550047	2031.B10.gz43_266223
M00055138A:E08	M00055137B:B11	ES 202	556630	2031.B14.gz43_266287
M00055138A:F01		ES 202	558477	2031.B16.gz43_266319
M00055138A-G08	M00055138A;E08	ES 202	558463	2031.C04.gz43_266128
M00055138B.H12	M00055138A:F01		558544	2031.C05.gz43_266144
M00055138C:A07		ES 202	558619	2031.C06.gz43_266160
M00055139B:E10		ES 202	446873	2031.C11.gz43_266240
M00055149B:F03	M00055138C:A07	ES 202	556328	2031.C12.gz43_266256
M00055140A;G07   ES 202   557893   2031,D22_gz43_26     M00055143B;C11   ES 202   555754   2031,G02_gz43_26     M00055144A;A11   ES 202   561245   2031,G02_gz43_26     M00055144A;E09   ES 202   450630   2031,G06_gz43_26     M00055144B;A12   ES 202   450630   2031,G06_gz43_26     M00055144B;E03   ES 202   490032   2031,G11_gz43_26     M00055146A;B12   ES 202   490032   2031,H18_gz43_26     M00055146A;B12   ES 202   211273   2031,104_gz43_26     M00055146A;B16   ES 202   289238   2031,J13_gz43_26     M00055149B;F09   ES 203   557783   2031,K05_gz43_26     M00055149B;F09   ES 203   5553600   2031,K05_gz43_26     M00055149B;C11   ES 203   5553600   2031,K05_gz43_26     M00055149B;C11   ES 203   5553600   2031,K05_gz43_26     M00055149B;C11   ES 203   5553600   2031,K05_gz43_26     M00055149B;C11   ES 203   5553600   2031,K05_gz43_26     M00055149B;C11   ES 203   5553600   2031,K05_gz43_26     M00055149B;C11   ES 203   5553600   2031,K05_gz43_26     M00055149B;C11   ES 203   5553600   2031,K05_gz43_26     M00055149B;C11   ES 203   5553600   2031,K05_gz43_26     M00055149B;C11   ES 203   5553600   2031,K05_gz43_26     M00055149B;C11   ES 203   5553600   2031,K05_gz43_26     M00055149B;C11   ES 203   5553600   2031,K05_gz43_26     M00055149B;C11   ES 203   5553600   2031,K05_gz43_26     M00055149B;C11   ES 203   5553600   2031,K05_gz43_26     M00055149B;C11   ES 203   2031,K05_gz43_26     M00055149B;C11   ES 203   2031,K05_gz43_26     M00055149B;C11   ES 203   2031,K05_gz43_26     M00055149B;C11   ES 203   2031,K05_gz43_26     M00055149B;C11   ES 203   2031,K05_gz43_26     M00055149B;C11   ES 203   2031,K05_gz43_26     M00055149B;C11   ES 203   2031,K05_gz43_26     M00055149B;C11   ES 203   2031,K05_gz43_26     M00055149B;C11   ES 203   2031,K05_gz43_26     M00055149B;C11   ES 203   2031,K05_gz43_26     M00055149B;C11   ES 203   2031,K05_gz43_26     M00055149B;C11   ES 203   2031,K05_gz43_26     M00055149B;C11   ES 203   2031,K05_gz43_26     M00055149B;C11   ES 203   2031,K05_gz43_26     M00055149B;C11   ES 203   20	M00055139B:E10	ES 202	553969	2031.D09.gz43_266209
M00055144B:A12	M00055139B:G03	ES 202	558230	2031.D11.gz43_266241
M00055144A:A11	M00055140A:G07	ES 202	557893	2031.D22.gz43_266417
M00055144A:E09   ES 202   450630   2031.G06.gz43_26   M00055144B:A12   ES 202   506372   2031.G11.gz43_26   M00055144B:F03   ES 202   490032   2031.H3 gz43_26   M00055146A:B12   ES 202   211273   2031.104.gz43_26   M00055146A:E09   ES 202   211273   2031.104.gz43_26   M00055146B:E09   ES 202   289328   2031.113.gz43_26   ES 202   289328   2031.J13.gz43_26   M00055149B:F09   ES 203   557783   2031.K05.gz43_26   M00055149B:F09   ES 203   553500   2031.K05.gz43_26   M00055149B:C11   ES 203   553500   2031.K05.gz43_26   M00055149B:C11   ES 203   553500   2031.K05.gz43_26   M00055149B:C11   ES 203   553500   2031.K05.gz43_26   M00055149B:C11   ES 203   553500   2031.K05.gz43_26   M00055149B:C11   ES 203   553500   2031.K05.gz43_26   M00055149B:C11   ES 203   553500   2031.K05.gz43_26   M00055149B:C11   ES 203   553500   2031.K05.gz43_26   M00055149B:C11   ES 203   553500   2031.K05.gz43_26   M00055149B:C11   ES 203   553500   2031.K05.gz43_26   M00055149B:C11   ES 203   553500   2031.K05.gz43_26   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 203   M00055149B:C11   ES 20	M00055143B:C11	ES 202	555754	2031.G02.gz43_266100
M00055144B:A12   ES 202   506372   2031.G11.gz43_26   M00055145B:F03   ES 202   490032   2031.H18.gz43_26   M00055146A:B12   ES 202   211273   2031.I04.gz43_26   M00055146B:E09   ES 202   562000   2031.I10.gz43_26   M00055148A:E06   ES 202   289328   2031.J13.gz43_26   M00055149B:F09   ES 203   557783   2031.K05.gz43_26   M00055149B:F09   ES 203   557500   2031.K05.gz43_26   M00055149C:A11   ES 203   553500   2031.K08.gz43_26   M00055149C:A11   ES 203   553500   2031.K08.gz43_26   M00055149C:A11   ES 203   553500   2031.K08.gz43_26   M00055149C:A11   ES 203   553500   2031.K08.gz43_26   M00055149C:A11   ES 203   553500   2031.K08.gz43_26   M00055149C:A11   ES 203   553500   2031.K08.gz43_26   M00055149C:A11   ES 203   553500   2031.K08.gz43_26   M00055149C:A11   ES 203   553500   2031.K08.gz43_26   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   ES 203   M00005149C:A11   M00005149C:A11   M00005149C:A11   M00005149C:A11   M00005149C:A11   M00005149C:A11   M00005149C:A11   M00005149C:A11   M00005149C:A11   M00005149C:A11   M00005149C:A11   M00005149C:A11   M00005149C:A11   M00005149C:A11   M00005149C:A11   M00005149C:A11	M00055144A:A11	ES 202	561245	2031.G05.gz43_266148
M00055145B:F03   ES 202   490032   2031.H18 g.rsl ³ 26   M00055146A:B12   ES 202   211273   2031.104 g.rsl ³ 26   M00055146B:E09   ES 202   56;2000   2031.H10 g.rsl ³ 26   M00055148A:E06   ES 202   289328   2031.J13 g.rsl ³ 26   M00055149B:F09   ES 203   557783   2031.K05 g.rsl ³ 26   M00055149C:A11   ES 203   553500   2031.K08 g.rsl ³ 26   M00055149C:A11   ES 203   553500   2031.K08 g.rsl ³ 26   M00055149C:A11   ES 203   553500   2031.K08 g.rsl ³ 26   M00055149C:A11   ES 203   553500   2031.K08 g.rsl ³ 26   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   ES 203   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00055149C:A11   M00051	M00055144A:E09	ES 202	450630	2031.G06.gz43_266164
M00055146A:B12         ES 202         211273         2031.04 gz43_26           M00055146B:E09         ES 202         562000         2031.110 gz43_26           M00055148A:E06         ES 202         289328         2031.J13 gz43_26           M00055149B:F09         ES 203         557783         2031.K05.gz43_26           M00055149C:A11         ES 203         553500         2031.K08.gz43_26	M00055144B:A12	ES 202	506372	2031.G11.gz43_266244
M00055146B:E09         ES 202         562000         2031.110.gx43_26           M00055148A:E06         ES 202         289328         2031.J13.gz43_26           M00055149B:F09         ES 203         557783         2031.K05.gz43_26           M00055149C:A11         ES 203         553500         2031.K08.gz43_26	M00055145B;F03	ES 202	490032	2031.H18.gz43_266357
M00055148A.E06 ES 202 289328 2031.I13.gz43 26 M00055149B:F09 ES 203 557783 2031.K05.gz43 26 M00055149C:A11 ES 203 553500 2031.K08.gz43 26	M00055146A:B12	ES 202	211273	2031.I04.gz43_266134
M00055149B:F09 ES 203 557783 2031.K05.gz43_26 M00055149C:A11 ES 203 553500 2031.K08.gz43_26	M00055146B;E09	ES 202	562000	2031.I10.gz43_266230
M00055149C:A11 ES 203 553500 2031.K08.gz43_26	M00055148A:E06	ES 202	289328	2031.J13.gz43_266279
M00055149C:A11 ES 203 553500 2031.K08.gz43_26				
	M00055149B:F09	ES 203	557783	2031.K05.gz43_266152
			553500	2031.K08.gz43_266200
				2031.L10.gz43_266233
	M00055151A:C10		557522	2031.L14.gz43_266297
M00055151A:F10 ES 203 610893 2031.L18.gz43_26	M00055151A:F10	ES 203	610893	2031.L18.gz43_266361

Table 13

Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055152A:B05	ES 203	490846	2031.M08.gz43_266202
M00055152A;E09	ES 203	549739	2031.M12.gz43_266266
M00055152B:A03	ES 203	557250	2031.M16.gz43_266330
M00055152D:C03	ES 203	468257	2031.N04.gz43_266139
M00055152D:H09	ES 203	497493	2031.N07.gz43_266187
M00055154D:F06	ES 203	549930	2031.O05.gz43_266156
M00055154D:G12	ES 203	524721	2031.O06.gz43_266172
M00055155D:B02	ES 203	557708	2031.O24.gz43_266460
M00055156A:D02	ES 203	561624	2031.P06.gz43_266173
M00055156B:C11	ES 203	558645	2031.P08.gz43_266205
M00055156C:A08	ES 203	558007	2031.P09.gz43_266221
M00055156C:D06	ES 203	561558	2031.P12.gz43_266269
M00055156D:A02	ES 203	561178	2031.P15.gz43_266317
M00055157A:B04	ES 203	556288	2031.P17.gz43_266349
M00055157C:C11	ES 203	559389	2032.A05.gz43 266526
M00055158A:D09	ES 203	555210	2032.A10.gz43 266606
M00055158D:C01	ES 203	394772	2032.A19.gz43 266750
M00055158D:D10	ES 203	560296	2032.A20.gz43_266766
M00055159B:B08	ES 203	561308	2032.B04.gz43 266511
M00055159B:G09	ES 203	561911	2032.B12.gz43_266639
M00055159C:B02	ES 203	215005	2032.B16.gz43_266703
M00055159D:F09	ES 203	554885	2032.B24.gz43 266831
M00055160C:D02	ES 203	553537	2032.C10.gz43 266608
M00055161A:C02	ES 203	561507	2032,C21,gz43 266784
M00055161A:E05	ES 203	558395	2032,C23.gz43_266816
M00055161B:A07	ES 203	452901	2032,D06,gz43 266545
M00055161D:H03	ES 203	447386	2032.D23.gz43 266817
M00055162A:B03	ES 203	562881	2032.E01.gz43_266466
M00055162A:C12	ES 203	451458	2032.E03.gz43 266498
M00055162A:G12	ES 203	561994	2032.E07.gz43 266562
M00055162B:B04	ES 203	561413	2032.E10.gz43 266610
M00055162C:E12	ES 203	553877	2032.E19.gz43_266754
M00055162C:G03	ES 203	323165	2032.E20.gz43 266770
M00055162D:B01	ES 203	555883	2032.E22.gz43 266802
M00055162D:G04	ES 203	561918	2032,E24.gz43 266834
M00055163B:F07	ES 203	561876	2032.F10.gz43 266611
M00055164C:C10	ES 203	558965	2032.G11.gz43 266628
M00055164D:C05	ES 203	555200	2032,G18.gz43 266740
M00055165A:F05	ES 203	559389	2032.G24.gz43 266836
M00055165B:G09	ES 203	559562	2032.H05.gz43 266533
M00055165C:H08	ES 203	558917	2032.H07.gz43 266565
M00055165D:C03	ES 203	553797	2032.H10.gz43 266613
M00055166B:D07	ES 203	452874	2032.H19.gz43_266757
M00055166C:G01	ES 203	488108	2032.H23.gz43_266821
MO0022TOOC:GOI	ES 203	400100	2032.F123.g243_200821

Table 13

A MOTO TO			
CloneID	ES No	ClusterID	SequenceName
M00055166D:F02	ES 203	559464	2032.I04.gz43_266518
M00055166D:F09	ES 203	555418	2032.I05.gz43_266534
M00055167A:A02	ES 203	493261	2032.I07.gz43_266566
M00055167B:A08	ES 203	557961	2032.I12.gz43 266646
M00055167B:H07	ES 203	415538	2032.I15.gz43_266694
M00055167D:B05	ES 203	558858	2032.I19.gz43_266758
M00055168B:F11	ES 203	557316	2032.J06.gz43_266551
M00055169B;F04	ES 203	559355	2032.J19.gz43_266759
M00055169B:H08	ES 203	550674	2032.J21.gz43_266791
M00055169D:A11	ES 203	558720	2032,K02.gz43_266488
M00055170A:A09	ES 203	557209	2032.K.06.gz43_266552
M00055170A:A11	ES 203	558755	2032.K07.gz43_266568
M00055170B:B06	ES 203	558022	2032.K12.gz43_266648
M00055170D:B09	ES 203	494198	2032.L01.gz43_266473
M00055171C:C01	ES 203	450566	2032.L23.gz43_266825
M00055172A:A01	ES 203	550085	2032.M02.gz43_266490
M00055172A:C09	ES 203	559015	2032.M05.gz43_266538
M00055172A:F03	ES 203	554604	2032.M06.gz43_266554
M00055172B:B04	ES 203	558900	2032.M09.gz43 266602
M00055172B:H07	ES 203	409262	2032.M12.gz43_266650
M00055172D:D04	ES 203	559057	2032.M15.gz43 266698
M00055172D:D07	ES 203	556542	2032.M16.gz43 266714
M00055172D:F12	ES 203	551693	2032.M19.gz43_266762
M00055174A:H12	ES 203	555202	2032.M24.gz43 266842
M00055174B:B04	ES 203	550175	2032.N01.gz43 266475
M00055176A:A02	ES 203	553047	2032.N11.gz43_266635
M00055176A:B03	ES 203	553713	2032.N13.gz43 266667
M00055176D:H01	ES 203	559752	2032,N24.gz43 266843
M00055177A:F05	ES 203	551475	2032.O01.gz43_266476
M00055177D:F07	ES 203	562989	2032.017.gz43_266732
M00055178A:C07	ES 203	558981	2032.O20.gz43 266780
M00055178A:D03	ES 203	559088	2032.022.gz43_266812
M00055179A:G08	ES 203	561975	2041.A02.gz43_266862
M00055179A:H11	ES 203	479851	2041.A03.gz43 266878
M00055179B:C07	ES 203	450289	2041.A04.gz43 266894
M00055179B:D05	ES 203	559050	2041.A05.gz43 266910
M00055179B:G07	ES 203	559000	2041.A08.gz43 266958
M00055179C:F11	ES 203	559383	2041.A10.gz43 266990
M00055179C:H02	ES 203	559710	2041.A11.gz43 267006
M00055179D:C10	ES 203	555739	2041.A14.gz43_267054
M00055181A:E01	ES 203	549829	2041.A20.gz43_267150
M00055181A:G02	ES 203	557644	2041.A22.gz43 267182
M00055181A:H01	ES 203	559194	2041.A24.gz43 267214
M00055181B:A10	ES 203	478511	2041.B01.gz43 266847

Table 13

Table 15			
CloneID	ES No	ClusterID	SequenceName
M00055181B:E06	ES 203	549472	2041.B03.gz43_266879
M00055181C:B07	ES 203	476199	2041.B07.gz43_266943
M00055181C:B12	ES 203	553002	2041.B08.gz43_266959
M00055181C:C09	ES 203	561513	2041.B09.gz43 266975
M00055181C:D06	ES 203	562569	2041.B11.gz43 267007
M00055181C:H01	ES 203	561351	2041.B17.gz43_267103
M00055181D:D05	ES 203	473742	2041.B21.gz43_267167
M00055182A:D07	ES 203	554520	2041.C05.gz43_266912
M00055182B:C07	ES 203	226324	2041.C08.gz43 266960
M00055182B:F05	ES 203	556632	2041.C09.gz43_266976
M00055182C:E09	ES 203	556881	2041.C17.gz43 267104
M00055182D:E06	ES 203	559333	2041.C21.gz43 267168
M00055182D:H08	ES 203	557961	2041.C23.gz43 267200
M00055183A:C06	ES 203	456517	2041.D01.gz43_266849
M00055183A:E10	ES 203	557714	2041.D02.gz43 266865
M00055183C:A02	ES 203	562840	2041.D08.gz43_266961
M00055183C:D07	ES 203	561963	2041.D12.gz43 267025
M00055184A:G02	ES 203	557935	2041.D19.gz43_267137
M00055184B:H01	ES 203	560308	2041.E02.gz43_266866
M00055184C:C07	ES 203	558212	2041.E04.gz43_266898
M00055184C:D02	ES 203	409612	2041.E05.gz43_266914
M00055184C:D11	ES 203	551879	2041.E07.gz43_266946
M00055184C:F01	ES 203	561513	2041.E08.gz43_266962
M00055185A:B01	ES 203	451761	2041.E12.gz43_267026
M00055185A:G11	ES 203	558582	2041.E14.gz43_267058
M00055185B:B01	ES 203	559883	2041.E16.gz43_267090
M00055185C:B01	ES 203	411113	2041.E22.gz43_267186
M00055185D:A02	ES 203	561687	2041.F03.gz43_266883
M00055185D:D11	ES 203	555394	2041.F06.gz43_266931
M00055185D:F07	ES 203	477046	20'41.F08.gz43_266963
M00055185D:H01	ES 203	502343	2041.F10.gz43_266995
M00055186A:D04	ES 203	552629	2041,F11.gz43_267011
M00055186A:E08	ES 203	557867	2041.F12.gz43_267027
M00055186C:A01	ES 203	459103	2041.F15.gz43_267075
M00055186C:A02	ES 203	481136	2041.F16.gz43_267091
M00055187A:F02	ES 203	516484	2041.G07.gz43_266948
M00055187A:F06	ES 203	492627	2041.G08.gz43_266964
M00055187A:G02	ES 203	552629	2041.G09.gz43_266980
M00055187B:C01	ES 203	364462	2041.G11.gz43_267012
M00055187C:C02	ES 203	551855	2041.G15.gz43_267076
M00055187C:E07	ES 203	549588	2041.G17.gz43_267108
M00055187D:A08	ES 203	557298	2041.G19.gz43_267140
M00055187D:F01	ES 203	560294	2041.G23.gz43_267204
M00055187D:G11	ES 203	557895	2041.H01.gz43_266853

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TABIC 15			
CloneID	ES No	ClusterID	SequenceName
M00055188A:G11	ES 203	559610	2041.H06.gz43_266933
M00055188A:H10	ES 203	187704	2041.H07.gz43_266949
M00055188B:E06	ES 203	559262	2041.H08.gz43_266965
M00055188B:G06	ES 203	555649	2041.H09.gz43_266981
M00055188C:A08	ES 203	448046	2041.H12.gz43_267029
M00055188C:F08	ES 203	549665	2041.H14.gz43_267061
M00055188D:D05	ES 203	559146	2041.H19.gz43_267141
M00055188D:F10	ES 203	561685	2041.H20.gz43_267157
M00055188D:H03	ES 203	559825	2041.H21.gz43_267173
M00055189A:C11	ES 203	558463	2041.H24.gz43_267221
M00055189B:B02	ES 203	452094	2041.I04.gz43_266902
M00055189B:B12	ES 203	560080	2041.I06.gz43_266934
M00055189C:C01	ES 203	413915	2041.I10.gz43_266998
M00055190A:A05	ES 203	523332	2041.I15.gz43_267078
M00055190A:F11	ES 203	455814	2041.I19.gz43_267142
M00055190C:G08	ES 203	560859	2041.I24.gz43_267222
M00055191B:A10	ES 203	488613	2041.J05.gz43_266919
M00055191B:E04	ES 203	560674	2041.J07.gz43_266951
M00055191C:C06	ES 203	447412	2041.J10.gz43_266999
M00055191D:C05	ES 203	560213	2041.J13.gz43_267047
M00055192A:A09	ES 203	559883	2041.J15.gz43_267079
M00055192C:A03	ES 203	499903	2041.J22.gz43_267191
M00055192C:D04	ES 203	491992	2041.K01.gz43_266856
M00055192C:E04	ES 203	550874	2041.K02.gz43_266872
M00055192C:H06	ES 203	558326	2041.K05.gz43_266920
M00055193A:A08	ES 203	560003	2041.K11.gz43_267016
M00055193A:C06	ES 203	451544	2041.K14.gz43_267064
M00055193B:A08	ES 203	624044	2041.K18.gz43_267128
M00055193C:C11	ES 203	539142	2041.L02.gz43_266873
M00055193C:E10	ES 203	504880	2041.L03.gz43_266889
M00055193D:G07	ES 203	552977	2041.L08.gz43_266969
M00055194A:A01	ES 203	495591	2041.L10.gz43_267001
M00055194A:E07	ES 203	558413	2041.L13.gz43_267049
M00055194B:C01	ES 203	559234	2041.L15.gz43_267081
M00055194B:G04	ES 203	513168	2041.L16.gz43_267097
M00055194D:C05	ES 203	549911	2041.M02.gz43_266874
M00055195A:B08	ES 203	549304	2041.M09.gz43_266986
M00055195A:C10	ES 203	550704	2041.M11.gz43_267018
M00055195A:E07	ES 203	143218	2041.M12.gz43_267034
M00055195B:B09	ES 203	450623	2041.M14.gz43_267066
M00055195B:C04	ES 203	614369	2041.M15.gz43_267082
M00055195C:F09	ES 203	561659	2041.N01.gz43_266859
M00055195C:H05	ES 203	561830	2041.N02.gz43_266875
M00055195D:B10	ES 203	555742	2041.N06.gz43_266939

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1 able 15			
CloneID	ES No	ClusterID	SequenceName
M00055195D:E11	ES 203	550652	2041.N09.gz43_266987
M00055196A;C04	ES 203	495074	2041.N12.gz43_267035
M00055196A:E01	ES 203	551977	2041.N13.gz43_267051
M00055196A:H07	ES 203	553237	2041.N15.gz43_267083
M00055196B:A09	ES 203	552879	2041,N17.gz43_267115
M00055196B:C06	ES 203	562336	2041.N18.gz43_267131
M00055196C:B07	ES 203	456697	2041.N24.gz43_267227
M00055196D:A10	ES 203	556538	2041.008.gz43_266972
M00055196D:F07	ES 203	481231	2041.O10.gz43_267004
M00055196D:H02	ES 203	562569	2041.O11.gz43_267020
M00055197B:A10	ES 203	559885	2041.O16.gz43_267100
M00055197B:B06	ES 204	558642	2041.O18.gz43_267132
M00055197C:D10	ES 204	625988	2041.O21.gz43_267180
M00055198A:E05	ES 204	460727	2041.P06.gz43_266941
M00055198B:H08	ES 204	490154	2041.P09.gz43_266989
M00055198C:C12	ES 204	551374	2041.P11.gz43_267021
M00055198C:F02	ES 204	557852	2041.P13.gz43_267053
M00055198C:G07	ES 204	561753	2041.P14.gz43_267069
M00055198D:A12	ES 204	551415	2041.P16.gz43_267101
M00055198D:B08	ES 204	556343	2041.P17.gz43_267117
M00055198D:G01	ES 204	550782	2041.P20.gz43_267165
M00055198D:G03	ES 204	491260	2041.P21.gz43_267181
M00055219B:B04	ES 204	554627	2043.A06.gz43_259273
M00055219C:C10	ES 204	556790	2043.A14.gz43_259401
M00055219C:H06	ES 204	558652	2043.A17.gz43_259449
M00055220B;H06	ES 204	490414	2043.B06.gz43_259274
M00055220D:G07	ES 204	557965	2043.B12.gz43_259370
M00055221A:D10	ES 204	549634	2043.B17.gz43_259450
M00055221D:A06	ES 204	558055	2043.C07,gz43_259291
M00055221D:H08	ES 204	467710	2043.C11.gz43_259355
M00055222A:A06	ES 204	460493	2043,C12.gz43_259371
M00055222A:C08	ES 204	464067	2043.C13.gz43_259387
M00055222B;A01	ES 204	552086	2043.C18.gz43_259467
M00055222B:A06	ES 204	451966	2043.C19.gz43_259483
M00055222D:B11	ES 204	558185	2043.D03.gz43_259228
M00055223B:C04	ES 204	552521	2043.D13.gz43_259388
M00055223D:F10	ES 204	621635	2043.D23.gz43_259548
M00055223D:H03	ES 204	559764	2043.D24.gz43_259564
M00055224B:E12	ES 204	561289	2043.E04.gz43_259245
M00055225B:H01	ES 204	560717	2043.E24.gz43_259565
M00055225C:E08	ES 204	550632	2043.F03.gz43_259230
M00055225D:G11	ES 204	491240	2043.F09.gz43_259326
M00055226B:F10	ES 204	556123	2043.F20.gz43_259502

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Table 13 CloneID	ES No	ClusterID	SequenceName
M00055226C:A11	ES 204	489275	2043.F23,gz43 259550
M00055226C:H05	ES 204	558679	2043.G08.gz43_259311
M00055227C:D02	ES 204	550815	2043,H09,gz43_259328
M00055227D:E07	ES 204	550571	2043.H16.gz43_259440
M00055227D:G10	ES 204	559101	2043.H17.gz43 259456
M00055228B:E07	ES 204	554181	2043.H21.gz43_259520
M00055229B:E12	ES 204	452598	2043.H21.gz43_239320 2043.H21.gz43_259377
M00055229B:H11	ES 204	559729	2043.I12.gz43_239377 2043.I15.gz43_259425
M00055229C:D01	ES 204	559053	2043.I17.gz43_259423 2043.I17.gz43_259457
M00055229D:D07	ES 204	543540	2043.I22.gz43_259537
M00055230A:H01	ES 204	551677	2043.J03.gz43 259234
M00055231D:G01	ES 204	560986	2043.J03.gz43_259538 2043.J22.gz43_259538
M00055232A:A09	ES 204	460680	2043.J22.gz43_259554 2043.J23.gz43_259554
M00055232C:D04	ES 204	557411	2043.K14.gz43 259411
M00055232C:F09	ES 204	559447	2043.L05.gz43 259268
M00055233D:G11	ES 204	559884	2043.L15.gz43_259428
M00055234A:G12	ES 204	552430	2043.L13.gz43_239428 2043.L18.gz43_259476
M00055235B:C01	ES 204	374281	2043.M12.gz43_259381
M00055236A:B01	ES 204	466887	2043,M19,gz43 259493
M00055236R:B01	ES 204	450193	2043.M23.gz43_259557
M00055236C:E05	ES 204	451720	2043.N03.gz43_259238
M00055236D:E04	ES 204 ES 204	558422	2043.N05.gz43_259286
M00055236D:E04	ES 204	557140	2043.N07.gz43 259302
M00055238D:D10	ES 204	557656	2043.N07.gz43_259559 2043.O23.gz43_259559
M00055239B:C05	ES 204	558230	2043.P05.gz43 259272
M00055239E;C05	ES 204	481362	2043.P10.gz43 259352
M00055239D:C01	ES 204	552019	2043.P10.gz43_239332 2043.P13.gz43_259400
M00055239D:C01	ES 204	557676	2043.P16.gz43_259448
M00055240A:B07	ES 204	562317	2043.P19.gz43 259496
M00055240B:E12	ES 204 ES 204	552249	
M00055240C:A03	ES 204	558093	2043.P24.gz43_259576 2044.A03.gz43_259613
M00055240C:A05	ES 204	43349	2044.A03.gz43_239613 2044.A04.gz43_259629
M00055240D:D12	ES 204	497119	2044.A04.gz43_239629 2044.A10.gz43 259725
M00055241A:C06	ES 204 ES 204	450025	2044,A10.gz43_259805
M00055241D:F10	ES 204	616134	2044,A13.gz43_239803 2044.B03.gz43_259614
M00055241D:F10 M00055242D:E02	ES 204 ES 204		
M00055243D:C06	ES 204	541784 486787	2044.B22.gz43_259918
M00055243D:C06 M00055244A:C06	ES 204 ES 204	238121	2044.C16.gz43_259823
M00055244A:C06 M00055244B:F02	ES 204 ES 204	558500	2044.C20.gz43_259887
	ES 204	38280	2044.D01.gz43_259584
M00055244B:G10			2044.D04.gz43_259632
M00055244D:C07 M00055244D:D09	ES 204 ES 204	560317 549493	2044.D09.gz43_259712
			2044.D10.gz43_259728
M00055244D:G08	ES 204 ES 204	549889	2044.D13.gz43_259776
M00055245B:H01	ES 204	629002	2044.D20.gz43_259888

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055245C:D04	ES 204	560507	2044.D22.gz43_259920
M00055245D:A04	ES 204	558355	2044.E03.gz43_259617
M00055245D:D01	ES 204	551305	2044.E06.gz43_259665
M00055246A:C01	ES 204	560317	2044.E09.gz43_259713
M00055246A:E10	ES 204	560669	2044.E10.gz43_259729
M00055246B:C04	ES 204	560204	2044.E14.gz43_259793
M00055246B:D10	ES 204	484086	2044.E15.gz43_259809
M00055246C:A10	ES 204	557389	2044.E16.gz43_259825
M00055246C:F08	ES 204	561579	2044.E17.gz43_259841
M00055247B:A11	ES 204	607430	2044.F04.gz43_259634
M00055247C:B01	ES 204	557389	2044.F05.gz43_259650
M00055248A:F10	ES 204	558461	2044.F14.gz43_259794
M00055248B:B03	ES 204	560099	2044.F15.gz43_259810
M00055248C:B11	ES 204	555340	2044.F22.gz43_259922
M00055248D:B12	ES 204	556829	2044.G02.gz43_259603
M00055249B:G08	ES 204	550830	2044.G18.gz43_259859
M00055249C:B12	ES 204	560059	2044.G22.gz43_259923
M00055250B:A05	ES 204	504415	2044.H06.gz43_259668
M00055250B:G09	ES 204	552357	2044.H09.gz43_259716
M00055250C:F03	ES 204	616196	2044.H10.gz43_259732
M00055251A:B09	ES 204	488143	2044.H18.gz43_259860
M00055251C:D01	ES 204	549052	2044.I06.gz43_259669
M00055251C:H07	ES 204	523732	2044.I11.gz43_259749
M00055251D:B09	ES 204	464205	2044.I14.gz43_259797
M00055251D:H11	ES 204	500737	2044.I18.gz43_259861
M00055252A:H09	ES 204	558670	2044.I24.gz43_259957
M00055252B:D07	ES 204	558785	2044.J03.gz43_259622
M00055252C:B04	ES 204	557256	2044.J05.gz43_259654
M00055252C:E02	ES 204	159419	2044.J08.gz43_259702
M00055252D:C10	ES 204	447380	2044.J14.gz43_259798
M00055253A:C07	ES 204	560205	2044.J17.gz43_259846
M00055253D:C03	ES 204	560261	2044.K01.gz43_259591
M00055254A:E02	ES 204	557928	2044.K05.gz43_259655
M00055254C:A06	ES 204	493359	2044.K11.gz43_259751
M00055254D:B10	ES 204	558861	2044.K15.gz43_259815
M00055255A:F11	ES 204	494130	2044.K24.gz43_259959
M00055255B:A10	ES 204	450405	2044.L01.gz43_259592
M00055255B:F05	ES 204	558512	2044.L04.gz43_259640
M00055256A:D12	ES 204	562256	2044.L16.gz43_259832
M00055256B:C07	ES 204	562256	2044.L21.gz43_259912
M00055256B:G05	ES 204	557928	2044.L24.gz43_259960
M00055257A:D01	ES 204	549456	2044.M16.gz43_259833
M00055257B:B10	ES 204	558134	2044.M17.gz43_259849
M00055257D:A11	ES 204	488574	2044.N04.gz43_259642

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055257D:E02	ES 204	554246	2044.N07.gz43_259690
M00055258A;C07	ES 204	423884	2044.N10.gz43_259738
M00055259A:G06	ES 204	138927	2044.006.gz43_259675
M00055259B:G11	ES 204	558559	2044.O14.gz43_259803
M00055259C:G10	ES 204	476373	2044.O17.gz43_259851
M00055260B:A05	ES 204	558098	2044,P04.gz43_259644
M00055260B:H06	ES 204	460049	2044,P08.gz43_259708
M00055260C:A11	ES 204	560144	2044.P09.gz43_259724
M00055260C:A12	ES 204	557734	2044.P10.gz43_259740
M00055260D:D04	ES 204	562926	2044.P20.gz43_259900
M00055261A:D11	ES 204	558357	2053.A03.gz43_260006
M00055261B:G12	ES 204	550863	2053.A12.gz43_260150
M00055261D:D09	ES 204	128962	2053.A20.gz43_260278
M00055262A:C05	ES 204	558281	2053.B02.gz43 259991
M00055262B:A11	ES 204	558120	2053,B05.gz43_260039
M00055262B:C01	ES 204	554373	2053.B06.gz43_260055
M00055262C:B08	ES 204	558052	2053,B11.gz43_260135
M00055262C:F05	ES 204	446557	2053.B14.gz43_260183
M00055263C:D02	ES 204	478087	2053.C10.gz43_260120
M00055263C:F10	ES 204	450242	2053.C13.gz43_260168
M00055263C:G09	ES 204	505858	2053.C15.gz43_260200
M00055263D:C03	ES 204	554000	2053.C19.gz43_260264
M00055264A:F03	ES 204	511276	2053.D05.gz43_260041
M00055264B:E06	ES 204	556019	2053.D08.gz43_260089
M00055264B:H12	ES 204	450507	2053.D10.gz43_260121
M00055264D:A03	ES 204	551272	2053.D14.gz43_260185
M00055264D:E09	ES 204	557713	2053.D18.gz43_260249
M00055265A:G01	ES 204	34381	2053.E02.gz43_259994
M00055265A:G07	ES 204	555564	2053.E03.gz43_260010
M00055265C:A04	ES 204	512721	2053.E10.gz43_260122
M00055265C:D09	ES 204	560520	2053.E13.gz43_260170
M00055265C:E01	ES 204	560678	2053.E14.gz43_260186
M00055265C:F01	ES 204	557561	2053.E15.gz43_260202
M00055266A:H08	ES 204	559910	2053.F03.gz43 260011
M00055266B:C12	ES 204	550074	2053.F05.gz43 260043
M00055266D:C09	ES 204	557561	2053.F14.gz43 260187
M00055267A:D01	ES 204	487961	2053.F21.gz43 260299
M00055267A:G11	ES 204	511847	2053.F22.gz43_260315
M00055267B:B06	ES 204	561666	2053.F24.gz43_260347
M00055267D:G08	ES 204	562269	2053.G18.gz43_260252
M00055268B:D12	ES 204	556447	2053.H02.gz43_259997
M00055268D:G09	ES 204	448450	2053.H15.gz43_260205
M00055270D:B05	ES 204	475578	2053.I15.gz43_260206
M00055271A:C05	ES 204	562992	2053.I19.gz43_260270
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CloneID	ES No	ClusterID	SequenceName
M00055271B:C01	ES 204	553665	2053.I24.gz43_260350
M00055271B:D02	ES 204	446964	2053.J01.gz43_259983
M00055272D:E04	ES 204	559321	2053.K01.gz43_259984
M00055274B:A10	ES 204	562314	2053.K21.gz43 260304
M00055274C:C06	ES 204	558867	2053.L05.gz43_260049
M00055274C;F02	ES 204	452506	2053.L10.gz43_260129
M00055274C:F10	ES 204	557115	2053.L11.gz43_260145
M00055274D:A11	ES 204	560957	2053.L12.gz43_260161
M00055274D:B10	ES 204	549052	2053.L15.gz43_260209
M00055274D:C10	ES 204	558263	2053.L18.gz43_260257
M00055275B:H06	ES 204	558360	2053.M03.gz43_260018
M00055275D:E12	ES 204	560621	2053.M12.gz43_260162
M00055275D:G09	ES 204	362109	2053.M13.gz43_260178
M00055275D:H08	ES 204	562871	2053.M15.gz43_260210
M00055276B;C09	ES 204	560278	2053.M20.gz43_260290
M00055277D:A02	ES 204	555564	2053.N05.gz43_260051
M00055279A:E03	ES 204	551798	2053.O05.gz43_260052
M00055279B:D02	ES 204	480960	2053.009.gz43_260116
M00055280A:C09	ES 204	560538	2053.O18.gz43 260260
M00055280C:G09	ES 204	560977	2053.P05.gz43_260053
M00055281A:E04	ES 204	558437	2053.P20.gz43_260293
M00055281A;F08	ES 204	558513	2053.P21.gz43_260309
M00055281B:D04	ES 204	481958	2053.P22.gz43_260325
M00055282A:A01	ES 204	559113	2054.A07.gz43_267326
M00055286A:H08	ES 204	559728	2054.C19.gz43_267520
M00055288B:D01	ES 204	559127	2054.E01.gz43_267234
M00055288B;D08	ES 204	399121	2054.E02.gz43_267250
M00055288C:A09	ES 205	549575	2054.E07.gz43_267330
M00055288D:A03	ES 205	558730	2054.E14.gz43_267442
M00055294B:C03	ES 205	484617	2054.H17.gz43_267493
M00055294B:D04	ES 205	559043	2054.H18.gz43_267509
M00055294B:G01	ES 205	559531	2054.H20.gz43_267541
M00055296A:C05	ES 205	561779	2054.I14.gz43_267446
M00055296C:E08	ES 205	558446	2054.I21.gz43_267558
M00055297A:C01	ES 205	558940	2054.I24.gz43_267606
M00055297D:C02	ES 205	500337	2054.J13.gz43_267431
M00055300A;B06	ES 205	551930	2054.K20.gz43_267544
M00055300C:F11	ES 205	559460	2054.L01.gz43 267241
M00055302B:B10	ES 205	624133	2054.M03.gz43_267274
M00055302B:F07	ES 205	561096	2054.M05.gz43_267306
M00055305C:D08	ES 205	560420	2054.O18.gz43_267516
M00055305D:F07	ES 205	448519	2054.O23.gz43_267596
M00055307B:G08	ES 205	451834	2054.P22.gz43_267581

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00055308D:C09	ES 205	551811	2055.A23.gz43_267966
M00055308D:E10	ES 205	559965	2055.B01.gz43_267615
M00055310B:E02	ES 205	138470	2055.B20.gz43_267919
M00055312A:D11	ES 205	559112	2055.C08.gz43_267728
M00055312A:E10	ES 205	440707	2055.C09.gz43_267744
M00055312B:F01	ES 205	559452	2055.C15.gz43_267840
M00055312D:A09	ES 205	558813	2055.C23.gz43_267968
M00055313D:E10	ES 205	560369	2055.D17.gz43_267873
M00055314B:F03	ES 205	430146	2055.D22.gz43_267953
M00055314B:G07	ES 205	551912	2055.D23.gz43_267969
M00055315C:A09	ES 205	617813	2055.E13.gz43_267810
M00055316B:B10	ES 205	477757	2055.E22.gz43_267954
M00055317C:D04	ES 205	555359	2055.F12.gz43_267795
M00055319B:H06	ES 205	559794	2055,G19.gz43_267908
M00055319C:C03	ES 205	553709	2055.G22.gz43_267956
M00055319C:C07	ES 205	552857	2055.G23.gz43_267972
M00055320A:F04	ES 205	553986	2055.H09.gz43_267749
M00055320D:E09	ES 205	558105	2055.H20.gz43_267925
M00055321A:A04	ES 205	555200	2055.I01.gz43_267622
M00055321A:D11	ES 205	558161	2055.I03.gz43_267654
M00055321B:B10	ES 205	558890	2055.I05.gz43_267686
M00055321D:C12	ES 205	559027	2055.I13.gz43_267814
M00055322A:C08	ES 205	558254	2055.I23.gz43_267974
M00055322C:G11	ES 205	559574	2055.J04.gz43_267671
M00055322D:A01	ES 205	552673	2055.J05.gz43_267687
M00055322D:C12	ES 205	446900	2055.J07.gz43_267719
M00055323D:A12	ES 205	562236	2055.K01.gz43_267624
M00055324C:H10	ES 205	448677	2055.K13.gz43_267816
M00055324D:B02	ES 205	555277	2055.K15.gz43_267848
M00055325A:E12	ES 205	473343	2055.K21.gz43_267944
M00055325A:H02	ES 205	554585	2055.K22.gz43_267960
M00055330C:F05	ES 205	451118	2055.M20.gz43_267930
M00055330D:C05	ES 205	558980	2055.M22.gz43_267962
M00055330D:H12	ES 205	559776	2055.N01.gz43_267627
M00055333C:F12	ES 205	451429	2055.N22.gz43_267963
M00055333C:H07	ES 205	555349	2055,N23.gz43_267979
M00055334C:E11	ES 205	560538	2055.O16.gz43 267868
M00055334D:G07	ES 205	554176	2055,O21.gz43_267948
M00055335A:B06	ES 205	497493	2055,O24.gz43_267996
M00055335D:A03	ES 205	472188	2055.P12.gz43_267805
M00055336D:B03	ES 205	558562	2056,A04,gz43_268046
M00055337D:B10	ES 205	558867	2056,A15.gz43_268222
M00055338A:A02	ES 205	453004	2056.A16.gz43_268238
M00055338A:A03	ES 205	559971	2056.A17.gz43_268254

Table 13

M00055338A;H10	A HIDTO XII			
M00055338B:F06	CloneID	ES No	ClusterID	SequenceName
M00055338C:F03	M00055338A:H10	ES 205	561116	2056.A21.gz43_268318
M00055338C:G04	M00055338B:F06	ES 205	560714	2056.A23.gz43_268350
M00055338C:H06	M00055338C:F03	ES 205	551912	2056.B04.gz43_268047
M00055338D:E11	M00055338C:G04	ES 205	552430	2056.B05.gz43_268063
M00055339A:B06	M00055338C:H06	ES 205	561024	2056.B07.gz43_268095
M00055340B:D10	M00055338D:E11	ES 205	512657	2056.B09.gz43_268127
M00055340B:C03	M00055339A:B06	ES 205	560091	2056.B10.gz43_268143
M00055340B:E06	M00055339B:D10	ES 205	562883	2056.B13.gz43_268191
M00055340C:D01	M00055340B:C03	ES 205	449405	2056.C06.gz43_268080
Noncosts340D:D01	M00055340B:E06	ES 205	560652	2056.C07.gz43_268096
M00055340D:F08	M00055340C:D01	ES 205	560369	2056.C09.gz43_268128
M00055341B:A01	M00055340D:D01	ES 205	560370	2056.C11.gz43_268160
M00055341B:C07	M00055340D:F08	ES 205	560717	2056.C13.gz43_268192
M00055341C:C01	M00055341B:A01	ES 205	558653	2056.C16.gz43_268240
M00055341C:F04		ES 205	560254	2056.C19.gz43_268288
M00055341C:G12	M00055341C;C01	ES 205	560252	2056.C24.gz43_268368
M00055341D:H07	M00055341C:F04	ES 205	627386	2056.D03.gz43 268033
NO0055342A:E08	M00055341C:G12	ES 205	552669	2056,D04.gz43 268049
M00055342B:C08	M00055341D:H07	ES 205	552357	2056.D08.gz43 268113
M00055342C:E03	M00055342A;E08	ES 205	562576	2056.D14.gz43_268209
M00055342C.H06	M00055342B:C08	ES 205	403419	2056.D19.gz43_268289
M00055342D:B02	M00055342C:E03	ES 205	496460	2056.D23.gz43_268353
M00055342D-F07	M00055342C:H06	ES 205	561124	2056.E03.gz43_268034
M00055342D:H05	M00055342D:B02	ES 205	163336	2056.E04.gz43_268050
M00055343A:C09	M00055342D:F07	ES 205	627139	2056.E08.gz43_268114
M00055343A-D08	M00055342D:H05	ES 205	561124	2056.E11.gz43_268162
M00055343A:G03	M00055343A:C09	ES 205	560199	2056.E13.gz43_268194
M00055343C:B11	M00055343A:D08	ES 205	560377	2056.E14.gz43_268210
M00055343C:G09   BS 205   560939   2056.E21.gx43 268   M00055343C:G10   ES 205   560939   2056.E22.gx43 268   M00055343D:C12   ES 205   560259   2056.F01.gx43 268   M00055343D:G13   IS 205   560932   2056.F01.gx43 268   M00055343D:G06   ES 205   560932   2056.F03.gx43 268   M00055343D:H04   ES 205   554101   2056.F06.gx43 268   M00055344D:H04   ES 205   559296   2056.F07.gx43 268   M00055344A:G11   ES 205   448285   2056.F12.gx43 268   M00055344C:C08   ES 205   618311   2056.F18.gx43 268   M00055344C:C08   ES 205   491240   2056.F18.gx43 268   M00055344C:C08   ES 205   557264   2056.F19.gx43 268   M00055344C:C09   ES 205   558890   2056.F12.gx43 268   M00055344C:C09   ES 205   558890   2056.F12.gx43 268   M00055344C:H09   ES 205   558890   2056.F24.gx43 268   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 205   ES 20	M00055343A:G03	ES 205	560868	2056.E16.gz43_268242
M00055343C:G10	M00055343C:B11	ES 205	400047	2056.E19.gz43_268290
M00055343D:C12	M00055343C:G09	ES 205	560939	2056.E21.gz43_268322
M00055343D:G03	M00055343C:G10	ES 205	560939	2056.E22.gz43_268338
M00055343D:G06	M00055343D:C12	ES 205	560259	2056.F01.gz43_268003
M00055344D:H04	M00055343D:G03	ES 205	560932	2056.F05.gz43_268067
M00055344A;G11	M00055343D:G06	ES 205	554101	2056.F06.gz43_268083
M00055344B:A12         ES 205         618311         2056.F13.gz43_266           M00055344C:C08         ES 205         491240         2056.F18.gz43_266           M00055344C:E04         ES 205         557264         2056.F19.gz43_266           M00055344C:F09         ES 205         558890         2056.F20.gz43_266           M00055344C:H09         ES 205         558806         2056.F24.gz43_266	M00055343D:H04	ES 205	559296	2056.F07.gz43_268099
M00055344C:C08         ES 205         491240         2056.F18.gz43 266           M00055344C:E04         ES 205         557264         2056.F19.gz43 266           M00055344C:F09         ES 205         558890         2056.F20.gz43 266           M00055344C:H09         ES 205         559806         2056.F24.gz43 266	M00055344A:G11	ES 205	448285	2056.F12.gz43_268179
M00055344C:E04         ES 205         557264         2056.F19.gz42 266           M00055344C:F09         ES 205         558890         2056.F20.gz43 266           M00055344C:H09         ES 205         559806         2056.F24.gz43 266	M00055344B:A12	ES 205	618311	2056.F13.gz43_268195
M00055344C:F09 ES 205 558890 2056.F20.gz43_268 M00055344C:H09 ES 205 559806 2056.F24.gz43_268	M00055344C:C08	ES 205	491240	2056.F18.gz43_268275
M00055344C:H09 ES 205 559806 2056.F24.gz43_268	M00055344C:E04	ES 205	557264	2056.F19.gz43_268291
	M00055344C:F09	ES 205	558890	2056.F20.gz43_268307
M00055244D-400 EC 205 550052 2056 C01 cm42 265	M00055344C:H09	ES 205	559806	2056.F24.gz43_268371
	M00055344D;A09	ES 205	550063	2056.G01.gz43_268004
		ES 205	559049	2056.G05.gz43_268068

Table 13

CloneID	ES No	ClusterID	SequenceName
M00055345B:B03	ES 205	558769	2056.G06.gz43_268084
M00055345B:F03	ES 205	559375	2056.G09.gz43_268132
M00055345D:A04	ES 205	551305	2056.G16.gz43_268244
M00055345D:D01	ES 205	559087	2056.G18.gz43_268276
M00055345D:D12	ES 205	559093	2056,G19.gz43_268292
M00055345D:E02	ES 205	559246	2056,G20,gz43_268308
M00055345D:E05	ES 205	558334	2056.G21.gz43_268324
M00055346A:B01	ES 205	560088	2056.G24.gz43_268372
M00055346A:E12	ES 205	418482	2056.H03.gz43_268037
M00055346B:G03	ES 205	509505	2056.H08.gz43_268117
M00055347A:C03	ES 205	559004	2056.H16.gz43_268245
M00055347C:F01	ES 205	559359	2056.I04.gz43_268054
M00055348B:A02	ES 205	559963	2056.I17.gz43_268262
M00055348B:B05	ES 205	103123	2056.I18.gz43_268278
M00055348B:F05	ES 205	552258	2056.I22.gz43_268342
M00055348B:H05	ES 205	550973	2056.I23.gz43_268358
M00055348D:A01	ES 205	559857	2056.J04.gz43_268055
M00055349A:F07	ES 205	549041	2056.J11.gz43_268167
M00055349C:G07	ES 205	458618	2056.J19.gz43_268295
M00055349C:H12	ES 205	562849	2056.J20.gz43_268311
M00055349D:F02	ES 205	556011	2056.J24.gz43_268375
M00055350A:F01	ES 205	556216	2056.K06.gz43_268088
M00055350B:B12	ES 205	449927	2056.K09.gz43_268136
M00055350B:D09	ES 205	560695	2056.K11.gz43_268168
M00055350C:G11	ES 205	553875	2056.K15.gz43_268232
M00055351A:A08	ES 205	553505	2056.K21.gz43_268328
M00055351A:C09	ES 205	560311	2056.K22.gz43_268344
M00055351B:D11	ES 205	560498	2056.L01.gz43_268009
M00055351B:H12	ES 205	512014	2056.L02.gz43_268025
M00055352B:E01	ES 205	555095	2056.L24.gz43_268377
M00055352B:E06	ES 205	560544	2056,M01.gz43_268010
M00055352B;H05	ES 205	557372	2056,M04,gz43_268058
M00055352C:A07	ES 205	553591	2056.M06.gz43_268090
M00055353B:B09	ES 205	455379	2056.M13.gz43_268202
M00055353C:A05	ES 205	426698	2056.M18.gz43_268282
M00055353D:C05	ES 205	558959	2056.M24.gz43_268378
M00055354A:A01	ES 205	558182	2056.N04.gz43_268059
M00055354A:G11	ES 205	471712	2056.N11.gz43_268171
M00055354A:H08	ES 205	561068	2056.N12.gz43_268187
M00055354C:C12	ES 205	238146	2056.N16.gz43_268251
M00055354C:E01	ES 205	184995	2056.N17.gz43_268267
M00055354C:F04	ES 205	560805	2056.N18.gz43_268283
M00055355A:A10	ES 205	487623	2056,N23,gz43_268363
M00055355A:H04	ES 205	562719	2056.O02.gz43_268028

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Table 13

CloneID	ES No	ClusterID	SequenceName
M00055355C:A11	ES 205	559910	2056.O07.gz43_268108
M00055356A;B06	ES 205	471272	2056.O13.gz43_268204
M00055356A:D04	ES 205	626061	2056.O14.gz43_268220
M00055356A;D09	ES 205	556365	2056.O15.gz43_268236
M00055356B:B04	ES 205	549781	2056.O17.gz43_268268
M00055356C:H02	ES 205	561108	2056.P01.gz43_268013
M00055356D:C11	ES 205	560174	2056.P07.gz43_268109
M00055356D;G09	ES 205	549464	2056.P08.gz43_268125
M00055357A;A09	ES 205	449751	2056.P10.gz43_268157
M00055357B;A08	ES 205	554591	2056.P15.gz43_268237
M00055357B:B01	ES 205	448989	2056.P16.gz43_268253
M00055357B:B07	ES 205	551412	2056.P17.gz43 268269
M00055357C:H07	ES 205	492821	2056.P22.gz43_268349
M00055358A:F09	ES 205	627297	2065.A06.gz43_268462
M00055358B:C01	ES 205	561485	2065.A09.gz43_268510
M00055358D:G04	ES 205	465610	2065.A17.gz43 268638
M00055359B:F03	ES 205	559495	2065.A21.gz43 268702
M00055359B:G09	ES 205	559676	2065.A23.gz43 268734
M00055359B:H07	ES 205	491635	2065,A24.gz43_268750
M00055359C:H09	ES 205	501534	2065.B08.gz43 268495
M00055359D:C12	ES 205	451124	2065.B11.gz43_268543
M00055359D:H02	ES 205	559828	2065.B15.gz43_268607
M00055360C;C05	ES 205	495799	2065.C02.gz43_268400
M00055360C:F08	ES 205	471718	2065.C05.gz43_268448
M00055360C:G11	ES 205	456469	2065.C06.gz43_268464
M00055360D:H10	ES 205	449593	2065,C12.gz43_268560
M00055361A;C01	ES 205	559022	2065.C15.gz43 268608
M00055361B:F12	ES 205	553615	2065.C20.gz43_268688
M00055361B:G08	ES 205	559675	2065.C21.gz43_268704
M00055361C:E05	ES 205	562459	2065.C22.gz43 268720
M00055361D:C05	ES 205	558927	2065.D02.gz43 268401
M00055361D:H12	ES 205	559696	2065.D05.gz43 268449
M00055362C:B06	ES 205	558412	2065.D11.gz43_268545
M00055362C:G08	ES 205	559675	2065,D14.gz43 268593
M00055362C:H07	ES 205	484091	2065,D15.gz43 268609
M00055363A:C03	ES 205	556947	2065.D22.gz43 268721
M00055363A;D02	ES 205	32021	2065.D24.gz43 268753
M00055363A;F07	ES 205	550562	2065.E02.gz43 268402
M00055363C;E02	ES 205	91178	2065,E08,gz43 268498
M00055364B:B01	ES 205	558504	2065.E15.gz43_268610
M00055364B;D01	ES 205	463304	2065,E17.gz43 268642
M00055364B;E10	ES 205	559963	2065,E18,gz43_268658
M00055364C:B08	ES 205	534054	2065,E20.gz43_268690
M00055364D:E09	ES 205	636651	2065.F03.gz43 268419

Table 13

CloneID	ES No	ClusterID	£N
Cioneid	ES NO	Clusterin	SequenceName
M00055365B:G10	ES 206	559919	2005 100 -42 208515
M00055365C:D12	ES 206	558334	2065.F09.gz43_268515
M00055365C:F11	ES 206	561825	2065.F11.gz43_268547 2065.F13.gz43_268579
M00055366A;B04	ES 206	556850	2065.F15.gz43_268611
M00055366A:H08	ES 206	551551	
M00055366B:C04	ES 206	511746	2065.F17.gz43_268643
M00055366C:B11	ES 206	505971	2065.F20.gz43_268691
M00055366C:B11	ES 206		2065.F23.gz43_268739
		527410	2065.G07.gz43_268484
M00055367A:B11	ES 206	556430	2065.G12.gz43_268564
M00055367D:A05	ES 206	523606	2065.H02.gz43_268405
M00055368C:G06	ES 206	482515	2065.H21.gz43_268709
M00055368D:E03	ES 206	558813	2065.H23.gz43_268741
M00055369A:H08	ES 206	556959	2065.I07.gz43_268486
M00055369C:D04	ES 206	557783	2065.I09.gz43_268518
M00055370A:B03	ES 206	335714	2065.I17.gz43_268646
M00055370B:F07	ES 206	549163	2065.J02.gz43_268407
M00055370C:B08	ES 206	562280	2065.J05.gz43_268455
M00055370C:D02	ES 206	552031	2065.J06.gz43_268471
M00055370D:F06	ES 206	549984	2065.J12.gz43_268567
M00055370D:H07	ES 206	560959	2065.J13.gz43_268583
M00055370D:H12	ES 206	554885	2065.J14.gz43_268599
M00055371A:B05	ES 206	508515	2065.J15.gz43_268615
M00055371A:H10	ES 206	562027	2065.J18.gz43_268663
M00055371B:D01	ES 206	561610	2065.J19.gz43_268679
M00055371B:F01	ES 206	553705	2065.J21.gz43_268711
M00055371D:B08	ES 206	484748	2065.K05.gz43_268456
M00055372A:H02	ES 206	552265	2065.K11.gz43_268552
M00055372B:B02	ES 206	556357	2065.K13.gz43_268584
M00055372B:E01	ES 206	632260	2065.K14.gz43_268600
M00055372B:F11	ES 206	561868	2065.K18.gz43_268664
M00055372C:E03	ES 206	553904	2065.K22.gz43_268728
M00055372D:C11	ES 206	556416	2065.L04.gz43_268441
M00055372D:G11	ES 206	516008	2065.L09.gz43_268521
M00055373B:A09	ES 206	490308	2065.L14.gz43_268601
M00055373C:F05	ES 206	466887	2065.M01.gz43_268394
M00055373C:H03	ES 206	446890	2065.M02.gz43_268410
M00055373C:H10	ES 206	543323	2065.M03.gz43_268426
M00055373D:B08	ES 206	498509	2065.M04.gz43_268442
M00055374A:B11	ES 206	555796	2065.M11.gz43_268554
M00055374A:E01	ES 206	636876	2065.M13.gz43_268586
M00055374B:D05	ES 206	562565	2065.M19.gz43_268682
M00055374B:F06	ES 206	559764	2065.M21.gz43_268714
M00055374C:F01	ES 206	376726	2065.N04.gz43_268443
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Table 13

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CloneID	ES No	ClusterID	SequenceName
M00055374D:C09	ES 206	504167	2065.N10.gz43 268539
M00055374D:F10	ES 206	558507	2065.N11.gz43_268555
M00055375B:H02	ES 206	482090	2065.N21.gz43 268715
M00055375B:H07	ES 206	555878	2065.N22.gz43 268731
M00055375C:C08	ES 206	448741	2065.N24.gz43 268763
M00055375C:F02	ES 206	143210	2065.O01.gz43_268396
M00055376A:A08	ES 206	561229	2065.O05.gz43_268460
M00055376B:A03	ES 206	474580	2065.O09.gz43_268524
M00055376B:A11	ES 206	561279	2065.O10.gz43_268540
M00055376B:B01	ES 206	453846	2065.O11.gz43_268556
M00055377A:B11	ES 206	207099	2065.O23.gz43_268748
M00055377B:E10	ES 206	562844	2065.P02.gz43_268413
M00055377C:G01	ES 206	461325	2065.P08.gz43_268509
M00055377C:H08	ES 206	450637	2065.P09.gz43_268525
M00055377D:F12	ES 206	562801	2065.P13.gz43_268589
M00055378A:B12	ES 206	235456	2065.P19.gz43_268685
M00055378D:B07	ES 206	559938	2066.A12.gz43_270570
M00055378D:D04	ES 206	562524	2066.A13.gz43_270586
M00055379A:D07	ES 206	468565	2066.A17.gz43_270650
M00055380A:C06	ES 206	448949	2066.B17.gz43_270651
M00055380D:H02	ES 206	562085	2066.C12.gz43_270572
M00055381A:F02	ES 206	549082	2066.C14.gz43_270604
M00055381B:C10	ES 206	562386	2066.C22.gz43_270732
M00055381C:G03	ES 206	562876	2066.D09.gz43_270525
M00055381D:D08	ES 206	560868	2066.D15.gz43_270621
M00055382B:E02	ES 206	493487	2066.D23.gz43_270749
M00055382C:D08	ES 206	558437	2066.E06.gz43_270478
M00055382C:H06	ES 206	446399	2066.E12.gz43_270574
M00055382D:C05	ES 206	562382	2066.E14.gz43_270606
M00055382D:D04	ES 206	459764	2066.E15.gz43_270622
M00055383A:F03	ES 206	562757	2066.E22.gz43_270734
M00055383B:H09	ES 206	551995	2066.F08.gz43_270511
M00055383C:A08	ES 206	562216	2066.F09.gz43_270527
M00055383D:F02	ES 206	498390	2066.F17.gz43_270655
M00055383D:G08	ES 206	454031	2066.F22.gz43_270735
M00055383D;H11	ES 206	559854	2066.F24.gz43_270767
M00055384A:F05	ES 206	562725	2066.G06.gz43_270480
M00055384B;D10	ES 206	562498	2066.G12.gz43_270576
M00055384C:G07	ES 206	562870	2066.G18.gz43_270672
M00055384D:H06	ES 206	550766	2066.G20.gz43_270704
M00055385C:C04	ES 206	549045	2066.H06.gz43_270481
M00055385C:F06	ES 206	561325	2066.H08.gz43_270513
M00055385C:G09	ES 206	450349	2066.H10.gz43_270545
M00055385D:D03	ES 206	418622	2066.H14.gz43_270609

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Table 13	PON	CI I TEL	
CloneID	ES No	ClusterID	SequenceName
M00055386A:G05	ES 206	562849	2066.H21.gz43_270721
M00055386D:G02	ES 206	558297	2066.I11.gz43_270562
M00055386D:H04	ES 206	557247	2066.I12.gz43_270578
M00055387B:H12	ES 206	550829	2066.I22.gz43_270738
M00055388A:A09	ES 206	553244	2066,J08.gz43_270515
M00055388A:B06	ES 206	560948	2066.J09.gz43_270531
M00055388B:B02	ES 206	465836	2066,J13,gz43_270595
M00055388B:E01	ES 206	636532	2066.J17.gz43_270659
M00055388B:H04	ES 206	557249	2066.J20.gz43_270707
M00055388C:D01	ES 206	635965	2066.J22.gz43_270739
M00055388D:A01	ES 206	555681	2066.J24.gz43_270771
M00055388D:F11	ES 206	556793	2066.K10.gz43_270548
M00055389A:D08	ES 206	238196	2066.K14.gz43_270612
M00055389C:F12	ES 206	282015	2066.K24.gz43_270772
M00055390A:A05	ES 206	515115	2066.L06.gz43_270485
M00055390C:E06	ES 206	50604	2066.L18.gz43_270677
M00055391B:D05	ES 206	499690	2066.M10.gz43_270550
M00055391B:D07	ES 206	453508	2066.M11.gz43_270566
M00055391B:H07	ES 206	418921	2066.M14.gz43_270614
M00055391B:H08	ES 206	555639	2066.M15.gz43_270630
M00055392A:H06	ES 206	446242	2066.N03.gz43_270439
M00055392C:G07	ES 206	551640	2066.N11.gz43_270567
M00055392D:A06	ES 206	562701	2066.N13.gz43_270599
M00055393B:F04	ES 206	44015	2066.N23.gz43_270759
M00055393C:B02	ES 206	549889	2066.N24.gz43_270775
M00055394B:C06	ES 206	561457	2066.O13.gz43_270600
M00055394B:D08	ES 206	556011	2066.O15.gz43_270632
M00055394D:F03	ES 206	554908	2066.P07.gz43_270505
M00055395A:C02	ES 206	446531	2066.P09.gz43_270537
M00055395A:E09	ES 206	452808	2066.P13.gz43_270601
M00055395B:C04	ES 206	561487	2066.P17.gz43_270665
M00055396B:C06	ES 206	560003	2067.A11.gz43_270938
M00055396B:G02	ES 206	561922	2067.A14.gz43_270986
M00055397A:B10	ES 206	562236	2067.A23.gz43_271130
M00055397A:C06	ES 206	154980	2067.A24.gz43_271146
M00055397A:H07	ES 206	556380	2067.B06.gz43_270859
M00055397B:F12	ES 206	562736	2067.B08.gz43_270891
M00055397D:A01	ES 206	562147	2067.B12.gz43_270955
M00055398A:C11	ES 206	561438	2067.B20.gz43_271083
M00055398B:A05	ES 206	469852	2067.B21.gz43_271099
M00055398B:C05	ES 206	456687	2067.B23.gz43_271131
M00055398C:A11	ES 206	555882	2067.B24.gz43_271147
M00055398C:D01	ES 206	448511	2067.C02.gz43_270796
M00055398C:F07	ES 206	511792	2067.C06.gz43_270860

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CloneID M00055399C:A08 M00055400B:H12	ES No ES 206	ClusterID 513715	SequenceName 2067.C21.gz43 271100
M00055400B:H12		513715	2067 (21 42 271100
	EC 200		
	ES 206	562029	2067.D19.gz43_271069
M00055400D:G01	ES 206	556382	2067.E04.gz43_270830
M00055401A:C08	ES 206	449258	2067.E06.gz43 270862
M00055401D:E03	ES 206	551731	2067.E21.gz43 271102
M00055402A:A05	ES 206	561206	2067.F02.gz43 270799
M00055402A:H03	ES 206	98869	2067.F06.gz43 270863
M00055402C:C12	ES 206	551928	2067.F22.gz43_271119
M00055402D:A11	ES 206	486834	2067.G02.gz43_270800
M00055402D:H04	ES 206	562059	2067.G06.gz43_270864
M00055403A:C07	ES 206	555856	2067.G11.gz43 270944
M00055403B:A05	ES 206	490060	2067.G16.gz43_271024
M00055403B:A10	ES 206	558180	2067,G18.gz43_271056
M00055403B:D04	ES 206	555726	2067.G21.gz43 271104
M00055403B:D07	ES 206	553173	2067.G22.gz43 271120
M00055403B:G09	ES 206	551675	2067.G23.gz43 271136
M00055403B:G12	ES 206	561236	2067.G24.gz43_271152
M00055404A:B10	ES 206	561325	2067.H10.gz43 270929
M00055404A:D08	ES 206	553766	2067.H12.gz43 270961
M00055404C:C11	ES 206	530715	2067.H23.gz43 271137
M00055405A:A02	ES 206	561236	2067.I05.gz43_270850
M00055405A:C01	ES 206.	452761	2067,I06.gz43 270866
M00055405A:G11	ES 206	419489	2067,I07.gz43 270882
M00055405B:H05	ES 206	554742	2067.I10.gz43 270930
M00055405B:H06	ES 206	473617	2067.I11.gz43 270946
M00055405C:C04	ES 206	63669	2067,I13.gz43_270978
M00055405C:H07	ES 206	559102	2067,I14.gz43 270994
M00055405D:G05	ES 206	551617	2067.I20.gz43_271090
M00055406B:E07	ES 206	562584	2067.J05.gz43_270851
M00055406B:F10	ES 206	562369	2067.J07.gz43_270883
M00055406D:A03	ES 206	491260	2067.J13.gz43 270979
M00055406D:C10	ES 206	487893	2067.J16.gz43_271027
M00055406D:G12	ES 206	557853	2067,J18,gz43 271059
M00055407A:F08 .	ES 206	465828	2067,J24.gz43 271155
M00055407B:G07	ES 206	550730	2067.K02.gz43 270804
M00055407C:E04	ES 206	465284	2067.K06.gz43 270868
M00055407C:G04	ES 206	556105	2067.K08.gz43 270900
M00055407C:G11	ES 206	560252	2067.K09.gz43 270916
M00055408A:A05	ES 206	561212	2067.K15.gz43 271012
M00055408A:F12	ES 206	66014	2067.K23.gz43 271140
M00055408A:G09	ES 206	446650	2067.L01.gz43 270789
M00055408B:G04	ES 206	447800	2067.L08.gz43 270901
M00055408C:C04	ES 206	451938	2067.L11.gz43_270949
M00055408C:E04	ES 206	559776	2067.L13.gz43_270981

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CloneID	ES No	ClusterID	SequenceName
M00055408D:E05	ES 206	548939	2067.L17.gz43_271045
M00055408D:E11	ES 206	558720	2067.L18.gz43_271061
M00055409A:E06	ES 206	100821	2067.L23.gz43_271141
M00055409A:E10	ES 206	516043	2067.L24.gz43_271157
M00055410C:G08	ES 206	550135	2067.N05.gz43_270855
M00055410C:H11	ES 206	562001	2067.N08.gz43_270903
M00055410D:B02	ES 206	288626	2067.N09.gz43_270919
M00055410D:G01	ES 206	633189	2067.N11.gz43_270951
M00055411A:H04	ES 206	558583	2067.N16.gz43_271031
M00055411B:D12	ES 206	356058	2067.N22.gz43_271127
M00055411C:E02	ES 206	450060	2067.O03.gz43_270824
M00055412A:C05	ES 206	440833	2067.O14.gz43_271000
M00055412A:F09	ES 206	556734	2067.O15.gz43_271016
M00055412B:A07	ES 206	495958	2067.O16.gz43_271032
M00055412D:A12	ES 206	560838	2067.P03.gz43_270825
M00055413A:A02	ES 206	540000	2067.P07.gz43_270889
M00055413C:B09	ES 206	446171	2067.P19.gz43_271081
M00042583D:F03	ES 207	452204	2078.A10.gz43_269168
M00042583D:F11	ES 207	506901	2078.A11.gz43_269184
M00042586B:A04	ES 207	451722	2078.A22.gz43_269360
M00042586B:A09	ES 207	451933	2078.A23.gz43_269376
M00042586B:A10	ES 207	448453	2078.A24.gz43_269392
M00042586C:E01	ES 207	451923	2078.B03.gz43_269057
M00042586D:C07	ES 207	451868	2078.B06.gz43_269105
M00042587B:G07	ES 207	452260	2078.B09.gz43_269153
M00042588A:G10	ES 207	508088	2078.B16.gz43_269265
M00042588C:E07	ES 207	505858	2078.B19.gz43_269313
M00042589A:G06	ES 207	452236	2078.B21.gz43_269345
M00042589B:D04	ES 207	504007	2078.B22.gz43_269361
M00042589B:E03	ES 207	452066	2078.B23.gz43_269377
M00042589D:D08	ES 207	415825	2078.C08.gz43_269138
M00042589D:F02	ES 207	452182	2078.C09.gz43_269154
M00042590B:G02	ES 207	507349	2078.C11.gz43_269186
M00042590C:C09	ES 207	418340	2078.C12.gz43_269202
M00042621B:G01	ES 207	452257	2078,C16.gz43_269266
M00042621D:A03	ES 207	501401	2078.C19.gz43_269314
M00042622C:H12	ES 207	452315	2078.D03.gz43_269059
M00042623B:B09	ES 207	451802	2078.D06.gz43_269107
M00042623D:C02	ES 207 .	447210	2078.D09.gz43_269155
M00042624D:B05	ES 207	452462	2078.D14.gz43_269235
M00042625B:H10	ES 207	517247	2078.D20.gz43_269331
M00042625D:C07	ES 207	511746	2078.E06.gz43_269108
M00042625D:E08	ES 207	513888	2078.E07.gz43_269124

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1 able 13			
CloneID	ES No	ClusterID	SequenceName
M00042626B:A12	ES 207	452432	2078.E10.gz43_269172
M00042626D:D12	ES 207	452031	2078.E16.gz43_269268
M00042627B;A10	ES 207	452376	2078.E18.gz43_269300
M00042628B;G11	ES 207	451401	2078.F04.gz43_269077
M00042628C:G10	ES 207	516522	2078.F06.gz43_269109
M00042628D:F12	ES 207	447904	2078,F08,gz43_269141
M00042629A:E11	ES 207	452052	2078.F11.gz43_269189
M00042630B:C04	ES 207	451885	2078.F20.gz43_269333
M00042630C:C12	ES 207	503499	2078.F22.gz43_269365
M00042951D:C05	ES 207	451841	2078.G05.gz43_269094
M00042952A:H12	ES 207	452324	2078.G10.gz43_269174
M00042952B:A08	ES 207	500853	2078.G11.gz43_269190
M00042952B:C08	ES 207	451887	2078.G12.gz43_269206
M00042952C:H09	ES 207	452323	2078.G16.gz43_269270
M00042953B:D02	ES 207	504560	2078.H01.gz43_269031
M00042954A:F04	ES 207	452142	2078.H12.gz43_269207
M00042954D:D04	ES 207	451993	2078.H18.gz43_269303
M00042955D:C02	ES 207	502815	2078.I02.gz43_269048
M00042955D:H03	ES 207	448594	2078.I04.gz43_269080
M00042956A:H02	ES 207	448090	2078,I05.gz43_269096
M00042957A:D06	ES 207	451994	2078.I12.gz43_269208
M00042957D:C09	ES 207	451929	2078.I18.gz43_269304
M00042958B:H04	ES 207	452981	2078.J02.gz43_269049
M00042958C:D04	ES 207	512721	2078,J03,gz43_269065
M00042958C:G10	ES 207	516018	2078.J04.gz43_269081
M00042959A;B07	ES 207	423578	2078.J06.gz43_269113
M00042959A:E08	ES 207	514142	2078.J08.gz43_269145
M00042959B;E11	ES 207	514160	2078.J11.gz43_269193
M00042959C;C06	ES 207	452615	2078,J12.gz43_269209
M00042959D:A05	ES 207	510169	2078.J14.gz43_269241
M00042960A:B10	ES 207	448332	2078.J17.gz43_269289
M00042960D:C11	ES 207	512051	2078.K03.gz43_269066
M00042962A:G04	ES 207	452899	2078.K12.gz43_269210
M00042962B:A03	ES 207	446438	2078.K13.gz43_269226
M00042962C:D05	ES 207	513155	2078.K16.gz43_269274
M00042962D:B09	ES 207	511351	2078.K18.gz43_269306
M00042963B:A02	ES 207	452459	2078.K23.gz43_269386
M00042963B:E12	ES 207	514594	2078.L02.gz43_269051
M00042963D:F11	ES 207	452801	2078.L07.gz43_269131
M00042964C:D06	ES 207	513156	2078.L16.gz43_269275
M00042964C:D10	ES 207	513168	2078.L17.gz43_269291
M00042964D:C02	ES 207	512059	2078.L21,gz43_269355
M00042965A:B03	ES 207	452506	2078.L23.gz43_269387
M00042965A:G02	ES 207	516415	2078,M01.gz43_269036

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1 able 13			
CloneID	ES No	ClusterID	SequenceName
M00042965B:G08	ES 207	452898	2078.M04.gz43_269084
M00042965C:A05	ES 207	452454	2078.M06.gz43_269116
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M00042967A:E06	ES 207	513585	2078.M18.gz43_269308
M00042969B:E01	ES 207	505679	2078.N04.gz43_269085
M00042969B:G10	ES 207	452279	2078.N05.gz43_269101
M00042970D:D02	ES 207	452695	2078.N16.gz43_269277
M00042971A:B09	ES 207	510717	2078.N17.gz43_269293
M00042971A:D12	ES 207	452662	2078.N19.gz43_269325
M00042972C:F07	ES 207	451618	2078.O04.gz43_269086
M00042972C:F09	ES 207	373239	2078.O05.gz43_269102
M00042973B:B10	ES 207	452500	2078.O09.gz43_269166
M00042974D:B07	ES 207	456545	2078.O17.gz43_269294
M00042975C:A08	ES 207	450805	2078.O21.gz43_269358
M00042975D:G07	ES 207	453124	2078.P01.gz43_269039
M00042976B:F05	ES 207	506920	2078.P03.gz43_269071
M00042978B:F03	ES 207	452833	2078.P17.gz43_269295
M00042978D:E02	ES 207	27534	2078.P20.gz43_269343
M00042982C:G02	ES 207	448687	2079.A18.gz43_271818
M00042983A:E09	ES 207	504880	2079.A22.gz43_271882
M00042983D;A03	ES 207	456756	2079.B06.gz43_271627
M00042985B:C05	ES 207	448200	2079.B14.gz43_271755
M00042985B:F05	ES 207	506372	2079.B15.gz43_271771
M00042985C:B03	ES 207	451811	2079.B16.gz43_271787
M00042985C:D02	ES 207	504501	2079.B17.gz43_271803
M00042985C:D04	ES 207	451491	2079.B18.gz43_271819
M00042986B:E06	ES 207	447445	2079.B22.gz43_271883
M00042986C:B07	ES 207	501534	2079.B24.gz43_271915
M00042987C:E04	ES 207	505226	2079.C06.gz43_271628
M00042988D:G10	ES 207	452220	2079.C13.gz43_271740
M00042989C:A06	ES 207	510254	2079.C15.gz43_271772
M00042989C:B07	ES 207	511348	2079.C16.gz43_271788
M00042989D:A07	ES 207	509505	2079.C20.gz43_271852
M00042990A:E05	ES 207	448813	2079.C21.gz43_271868
M00042990B:A08	ES 207	500795	2079.C23.gz43 271900
M00042991B:G01	ES 207	452874	2079.D05.gz43_271613
M00042991C:C02	ES 207	238196	2079.D06.gz43_271629
M00042991D:C09	ES 207	452603	2079.D09.gz43_271677
M00042991D:F06	ES 207	452830	2079.D12.gz43 271725
M00042992B:B03	ES 207	451812	2079.D15.gz43_271773
M00042992B:B11	ES 207	502343	2079.D16.gz43_271789
M00042992D:H02	ES 207	75212	2079.D19.gz43_271837
M00042993A:D12	ES 207	513178	2079.D21.gz43_271869
M00042993B:C12	ES 207	446520	2079.D23.gz43 271901

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M00042996D:D12	ES 207	453804	2079.E21.gz43_271870
M00042997B:B07	ES 207	451806	2079.E23.gz43_271902
M00042999A:D01	ES 207	403634	2079.F07.gz43_271647
M00043000C:H06	ES 207	509027	2079.F14.gz43_271759
M00055413C:G09	ES 207	556126	2067.P21.gz43_271113
M00055414C:C02	ES 207	456520	2068.A13.gz43_271354
M00055414D:G10	ES 207	482425	2068.A21.gz43_271482
M00055415D:C12	ES 207	561500	2068.B15.gz43_271387
M00055415D:E10	ES 207	204557	2068.B17.gz43_271419
M00055416A:C04	ES 207	561497	2068.B21.gz43_271483
M00055416B:F11	ES 207	561770	2068.C07.gz43_271260
M00055416D:B03	ES 207	562477	2068.C12.gz43_271340
M00055416D:D11	ES 207	558679	2068.C14.gz43_271372
M00055416D:H11	ES 207	555880	2068.C18.gz43_271436
M00055417A:G11	ES 207	131130	2068.C21.gz43_271484
M00055417C:A05	ES 207	550166	2068.D04.gz43_271213
M00055417C:G03	ES 207	561112	2068.D05.gz43_271229
M00055417D:D08	ES 207	557760	2068.D08.gz43_271277
M00055417D:H08	ES 207	470667	2068.D10.gz43_271309
M00055418A:C12	ES 207	561454	2068.D12.gz43_271341
M00055418A:E07	ES 207	561682	2068.D15.gz43_271389
M00055418A:F03	ES 207	560252	2068.D16.gz43_271405
M00055418A:H04	ES 207	554774	2068.D17.gz43_271421
M00055418C:D08	ES 207	418562	2068.E05.gz43_271230
M00055418D:A03	ES 207	552561	2068.E10.gz43_271310
M00055418D:B05	ES 207	630348	2068.E11.gz43_271326
M00055419D:D04	ES 207	561646	2068.F04.gz43_271215
M00055419D:H07	ES 207	550018	2068.F09.gz43_271295
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M00055422D:E12	ES 207	631966	2068.I12.gz43_271346
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M00055423A:D02	ES 207	560118	2068.I15.gz43_271394
M00055423A:G08	ES 207	449356	2068.I16.gz43_271410

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Table 13

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M00055425B:D12	M00055424B:D04	ES 207	559380	2068.K01.gz43_271172
M00055425B:F06	M00055425A:H03	ES 207	554953	2068.K14.gz43_271380
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Table 13

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Table 13

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M00043086D:B09         ES 208         453248         2089.F08.gz43_269825           M00043087B:D10         ES 208         449430         2089.F13.gz43_269905           M00043087B:D07         ES 208         395536         2089.F16.gz43_269953           M00043089A:D06         ES 208         520616         2089.F21.gz43_270033           M0004309DB:H06         ES 208         524470         2089.F24.gz43_270081				
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M00043088B:D07         ES 208         395536         2089.F16.gz43_269953           M00043089A:D06         ES 208         520616         2089.F21.gz43_270033           M00043099B:H06         ES 208         524470         2089.F24.gz43_270081				
M00043089A:D06         ES 208         520616         2089.F21.gz43_270033           M00043090B:H06         ES 208         524470         2089.F24.gz43_270081			117 10 1	
M00043090B:H06 ES 208 524470 2089.F24.gz43_270081				
M00043090D:H07 ES 208 524478 2089.G03.gz43_269746				
	M00043090D:H07	ES 208	524478	2089.G03.gz43_269746

Table 13

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CloneID	ES No	ClusterID	SequenceName
M00043091C:H05	ES 208	530656	2089.G07.gz43_269810
M00043091C:H08	ES 208	454910	2089.G08.gz43_269826
M00043091D:C01	ES 208	454374	2089.G09.gz43_269842
M00043091D:F03	ES 208	402242	2089.G11.gz43_269874
M00043092D:D06	ES 208	454531	2089.G18.gz43_269986
M00043093A:A06	ES 208	454177	2089.G20.gz43 270018
M00043093D:F12	ES 208	454701	2089.H02.gz43 269731
M00043094A:F01	ES 208	529219	2089.H03.gz43 269747
M00043094C:A12	ES 208	525023	2089.H06.gz43 269795
M00043096A:B02	ES 208	449202	2089.H12.gz43 269891
M00043096A:E01	ES 208	528134	2089.H13.gz43 269907
M00043096C:D02	ES 208	454509	2089.H16.gz43_269955
M00043096C:H04	ES 208	451972	2089.H17.gz43 269971
M00043097A:D11	ES 208	527873	2089.H21.gz43_270035
M00043097A:F06	ES 208	387530	2089.H22.gz43_270051
M00043097D:B12	ES 208	454226	2089.I02.gz43_269732
M00043100C:D08	ES 208	453533	2089.I15.gz43_269940
M00043100D:C12	ES 208	520057	2089.I17.gz43_269972
M00043101A:F12	ES 208	522548	2089.I20.gz43 270020
M00043101C:F12	ES 208	453846	2089.I22.gz43 270052
M00043102A:B10	ES 208	453364	2089.J01.gz43_269717
M00043102A:G12	ES 208	523590	2089.J04.gz43_269765
M00043102B:F05	ES 208	453761	2089.J05.gz43_269781
M00043102D:C05	ES 208	446728	2089.J09.gz43 269845
M00043102D:F11	ES 208	453766	2089.J11.gz43_269877
M00043103A:G05	ES 208	449335	2089.J12.gz43_269893
M00043104B:C09	ES 208	453494	2089.J20.gz43_270021
M00043105A.F02	ES 208	453809	2089.K01.gz43_269718
M00043106B:F07	ES 208	450287	2089.K11.gz43 269878
M00043106C:D05	ES 208	453572	2089.K12.gz43_269894
M00043107A:E07	ES 208	453726	2089.K17.gz43_269974
M00043107D:H04	ES 208	450566	2089.K21.gz43_270038
M00043108B:A01	ES 208	453132	2089.K23.gz43_270070
M00043108B:D12	ES 208	453549	2089.L01.gz43 269719
M00043109C:F04	ES 208	454720	2089.L06.gz43 269799
M00043131B:A11	ES 208	454134	2089.L07.gz43 269815
M00043131B:G10	ES 208	454825	2089.L11.gz43 269879
M00043131C:A11	ES 208	451391	2089.L13.gz43_269911
M00043131D:B02	ES 208	525719	2089.L15.gz43_269943
M00043132C:D02	ES 208	454483	2089.L22.gz43_270055
M00043134C:D06	ES 208	454518	2089.M09.gz43_269848
M00043135C:E07	ES 208	454550	2089,M12.gz43_269896
M00043135D:A11	ES 208	454126	2089.M13.gz43_269912
M00043135D:C07	ES 208	526575	2089.M14.gz43 269928

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1 able 15			
CloneID	ES No	ClusterID	SequenceName
M00043136A:D03	ES 208	527361	2089.M17.gz43_269976
M00043137C:D02	ES 208	452936	2089.N04.gz43_269769
M00043137D:D10	ES 208	454438	2089.N06.gz43_269801
M00043138A:H03	ES 208	523753	2089.N09.gz43_269849
M00043138B:B08	ES 208	407275	2089.N10.gz43_269865
M00043138B:F02	ES 208	93125	2089.N12.gz43_269897
M00043138B:G11	ES 208	437064	2089.N13.gz43 269913
M00043138C;D09	ES 208	449394	2089.N15.gz43_269945
M00043139C;A05	ES 208	524622	2089.N23.gz43_270073
M00043139D:A06	ES 208	524624	2089.O02.gz43_269738
M00043140C:D03	ES 208	527446	2089.O04.gz43_269770
M00043140C:H11	ES 208	530919	2089.O07.gz43_269818
M00043141C:C12	ES 208	415326	2089.O10.gz43_269866
M00043141D:A12	ES 208	454129	2089.O13.gz43_269914
M00043144D:H01	ES 208	454050	2089.P03.gz43_269755
M00043146C:D10	ES 208	520595	2089.P09.gz43_269851
M00043146C:F10	ES 208	453756	2089.P10.gz43_269867
M00043146D:H03	ES 208	455941	2089.P13.gz43_269915
M00043147D:H03	ES 208	524363	2089.P15.gz43_269947
M00043148C:E01	ES 208	453692	2089.P17.gz43_269979
M00043149B:A01	ES 208	453202	2089.P20.gz43_270027
M00043149D:F02	ES 208	522220	2090.A03.gz43_273939
M00043150A:B12	ES 208	518949	2090.A04.gz43_273955
M00043151B:D02	ES 208	454460	2090.A12.gz43_274083
M00043152A:F10	ES 208	454731	2090.A17.gz43_274163
M00043152A:G08	ES 208	456723	2090.A18.gz43_274179
M00043152B:H04	ES 208	530939	2090.A23.gz43_274259
M00043153B:B09	ES 208	519378	2090.B01.gz43_273908
M00043156C:E05	ES 208	528404	2090.C03.gz43_273941
M00043158A:F03	ES 208	522869	2090.C12.gz43_274085
M00043159A:C01	ES 208	526449	2090.C18.gz43_274181
M00043160B:E05	ES 208	521840	2090.D02.gz43_273926
M00043162D:E06	ES 208	453730	2090.D17.gz43_274166
M00043168C:F03	ES 208	446768	2090.F05.gz43_273976
M00043169A:F05	ES 208	453804	2090.F09.gz43 274040
M00043169A:H08	ES 208	524100	2090.F11.gz43_274072
M00043169C:D10	ES 208	453569	2090.F15.gz43_274136
M00043171B:G05	ES 208	523602	2090.G12.gz43_274089
M00043171D:G08	ES 208	450784	2090.G17.gz43_274169
M00043172A:D08	ES 208	401160	2090.G18.gz43_274185
M00043172C:B05	ES 208	234270	2090.G23.gz43_274265
M00043172C:G06	ES 208	453946	2090.H01.gz43_273914
M00043172D:H03	ES 208	128749	2090.H06.gz43_273994
M00043173B:F03	ES 208	453354	2090.H11.gz43_274074

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Table 15			
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M00043173C:B05	ES 208	453256	2090.H13.gz43_274106
M00043173C:D12	ES 208	520599	2090.H15.gz43_274138
M00043174A:G04	ES 208	454810	2090.H17.gz43_274170
M00043174B:B06	ES 208	413693	2090.H18.gz43_274186
M00043174C:C06	ES 208	526733	2090.H20.gz43_274218
M00043176C:A01	ES 208	525315	2090.I11.gz43_274075
M00043176C:E12	ES 208	454664	2090.I13.gz43_274107
M00042592A:H02	ES 209	539142	2092.A07.gz43_274486
M00042594C:D02	ES 209	455405	2092.A22.gz43_274726
M00042595A:B07	ES 209	447278	2092.B01.gz43_274391
M00042599D:E07	ES 209	455460	2092.B15.gz43_274615
M00042600D:B08	ES 209	455117	2092.B20.gz43_274695
M00042600D:E11	ES 209	452392	2092.B21.gz43_274711
M00042637C:E12	ES 209	536171	2092.B22.gz43_274727
M00042637D:B12	ES 209	455814	2092.B23.gz43_274743
M00042638D:G10	ES 209	455784	2092.C02.gz43_274408
M00042639D:H04	ES 209	455880	2092.C09.gz43_274520
M00042640C:C10	ES 209	455201	2092.C11.gz43_274552
M00042640D:D01	ES 209	455375	2092.C12.gz43_274568
M00043178C:C01	ES 209	454324	2090.I22.gz43_274251
M00043178C:G06	ES 209	529742	2090.I24.gz43 274283
M00043180C:B02	ES 209	452325	2090.J08.gz43_274028
M00043181C:C10	ES 209	526539	2090.J14.gz43_274124
M00043181C:F06	ES 209	529037	2090.J15.gz43_274140
M00043182B:C02	ES 209	454311	2090.J18.gz43_274188
M00043183A:C04	ES 209	454363	2090.K02.gz43_273933
M00043183C:B08	ES 209	525781	2090.K06.gz43 273997
M00043184D:A02	ES 209	447536	2090.K17.gz43 274173
M00043184D:G06	ES 209	432159	2090.K20.gz43_274221
M00043185B:D02	ES 209	527679	2090.L01.gz43_273918
M00043186C:A11	ES 209	524721	2090.L07.gz43_274014
M00043186D:A06	ES 209	524706	2090.L09.gz43_274046
M00043186D:B09	ES 209	425455	2090.L10.gz43_274062
M00043187B:G03	ES 209	530094	2090.L12.gz43 274094
M00043187C:E06	ES 209	528369	2090.L13.gz43_274110
M00043187D:H04	ES 209	530971	2090.L16.gz43 274158
M00043188D:B07	ES 209	525456	2090.L17.gz43 274174
M00043189C:F08	ES 209	528981	2090.M02.gz43_273935
M00043189D:B11	ES 209	452231	2090.M05.gz43_273983
M00043190A:B10	ES 209	454202	2090.M07.gz43_274015
M00043191C:G05	ES 209	404081	2090.M17.gz43_274175
M00043192B:H06	ES 209	453983	2090.M20.gz43 274223
M00043193C:G11	ES 209	523674	2090.N04.gz43_273968

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Table 15			
CloneID	ES No	ClusterID	SequenceName
M00043193C:H09	ES 209	452775	2090.N06.gz43_274000
M00043194A:G06	ES 209	453890	2090.N11.gz43_274080
M00043194C:A05	ES 209	453177	2090.N14.gz43_274128
M00043194D:E10	ES 209	289316	2090.N16.gz43_274160
M00043195D:A05	ES 209	518172	2090.N20.gz43_274224
M00043197D:H12	ES 209	454007	2090.O09.gz43_274049
M00043198A:F01	ES 209	453818	2090.O11.gz43_274081
M00043199A:F03	ES 209	452182	2090.O14.gz43_274129
M00043201D:D03	ES 209	454463	2090.P01.gz43_273922
M00043203B:E07	ES 209	528616	2090.P10.gz43_274066
M00043210C:E08	ES 209	454629	2090.P22.gz43_274258
M00043417A:H12	ES 209	539804	2092.C18.gz43_274664
M00043417B:F12	ES 209	455601	2092.C20.gz43_274696
M00043417C:F12	ES 209	447380	2092.C21.gz43_274712
M00043418A:A06	ES 209	422223	2092.C23.gz43_274744
M00043420C:C02	ES 209	455289	2092.D09.gz43_274521
M00043421C:G07	ES 209	449067	2092.D12.gz43_274569
M00043422B:C03	ES 209	455162	2092.D13.gz43_274585
M00043425C:H11	ES 209	539452	2092.E07.gz43_274490
M00043427C:E11	ES 209	544355	2092.E10.gz43_274538
M00043428C:A08	ES 209	531461	2092.E15.gz43_274618
M00043428D:C10	ES 209	455220	. 2092.E17.gz43_274650
M00043428D:E02	ES 209	536038	2092.E18.gz43_274666
M00043430D:C01	ES 209	455254	2092.E20.gz43_274698
M00043433A:F02	ES 209	537451	2092.F08.gz43_274507
M00043434A:H02	ES 209	452128	2092.F10.gz43_274539
M00043434B:D06	ES 209	535129	2092.F11.gz43_274555
M00043434B:E04	ES 209	452801	2092.F12.gz43_274571
M00043438B:E10	ES 209	404461	2092.G04.gz43_274444
M00043438C:D12	ES 209	535436	2092.G05.gz43_274460
M00043439C:D05	ES 209	535123	2092.G08.gz43_274508
M00043441D:A09	ES 209	455132	2092.G17.gz43_274652
M00043442A:D02	ES 209	534519	2092.G18.gz43_274668
M00043446B:H01	ES 209	456020	2092,H09.gz43_274525
M00043446C:E04	ES 209	455808	2092.H11.gz43_274557
M00043446C:E08	ES 209	400258	2092.H12.gz43_274573
M00043446C:E10	ES 209	456530	2092.H13.gz43_274589
M00043446C:G09	ES 209	547652	2092.H14.gz43_274605
M00043447A:F02	ES 209	455572	2092.H15.gz43_274621
M00043447C:A06	ES 209	454994	2092,H18.gz43_274669
M00043448B:E08	ES 209	455501	2092.H22.gz43_274733
M00043448C:D05	ES 209	456742	2092,H23.gz43_274749
M00043448C:G07	ES 209	538582	2092,H24.gz43_274765
M00043448D:H09	ES 209	451023	2092.I04.gz43_274446

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Table 13 CloneID	ES No	ClusterID	SequenceName
M00043449C:H09	ES 209	448946	2092.I09.gz43 274526
M00043449D:A06	ES 209	447534	2092.I10.gz43_274542
M00043449D:E09	ES 209	215366	2092.I11.gz43 274558
M00043450B:C11	ES 209	456249	2092.I14.gz43 274606
M00043450B:H05	ES 209	456783	2092.I15.gz43_274622
M00043450C;F11	ES 209	454563	2092.I17.gz43 274654
M00043451A;B02	ES 209	456103	2092.I20.gz43 274702
M00043451B:D08	ES 209	448250	2092.I21.gz43 274718
M00043451C:H03	ES 209	447238	2092.I24.gz43_274766
M00043452D:D05	ES 209	452830	2092,J08.gz43 274511
M00043453B:C06	ES 209	456254	2092.J11.gz43 274559
M00043453B:F02	ES 209	546121	2092.J12.gz43 274575
M00043453C:A06	ES 209	446866	2092.J13.gz43 274591
M00043453D:D02	ES 209	543855	2092.J19.gz43 274687
M00043455B:G11	ES 209	546838	2092.K04.gz43 274448
M00043455C:G07	ES 209	450914	2092.K07.gz43 274496
M00043457C:B12	ES 209	454621	2092.K12.gz43 274576
M00043459A:B08	ES 209	541901	2092.K17.gz43 274656
M00043460D:C03	ES 209	534054	2092.L08.gz43 274513
M00043460D:H01	ES 209	539353	2092.L09.gz43 274529
M00043462A:H06	ES 209	455834	2092.L14.gz43_274609
M00043462C:E12	ES 209	536225	2092.L17.gz43 274657
M00043462D:C09	ES 209	415326	2092.L19.gz43 274689
M00043465B:G08	ES 209	162851	2092.M08.gz43_274514
M00043468C:D08	ES 209	455379	2092.M17.gz43_274658
M00043470C:A01	ES 209	456001	2092.N05.gz43_274467
M00043472A;E06	ES 209	536415	2092.N11.gz43_274563
M00043473C:B11	ES 209	37186	2092.N15.gz43_274627
M00043473D:D11	ES 209	452618	2092.N18.gz43_274675
M00043474C:H05	ES 209	455864	2092.N20.gz43_274707
M00043475C:G08	ES 209	446866	2092.N23.gz43_274755
M00043478B:D01	ES 209	450724	2092.O06.gz43_274484
M00043481A:B01	ES 209	532904	2092.O14.gz43_274612
M00043481A:G02	ES 209	402534	2092.O16.gz43_274644
M00043483B:G01	ES 209	455808	2092.O22.gz43_274740
M00043483B:G11	ES 209	455814	2092.O23.gz43_274756
M00043484C:E12	ES 209	455492	2092.P05.gz43_274469
M00043484D:H08	ES 209	455855	2092.P07.gz43_274501
M00043485A:A04	ES 209	452910	2092.P08.gz43_274517
M00043485A;C04	ES 209	450658	2092.P09.gz43_274533
M00043485B:C09	ES 209	455256	2092.P10.gz43_274549
M00043486B:D02	ES 209	543429	2092.P13.gz43_274597
M00043486C:D02	ES 209	543431	2092.P15.gz43_274629
M00043486C:F07	ES 209	456567	2092.P17.gz43_274661

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Table 13			
CloneID	ES No	ClusterID	SequenceName
M00043489C:H06	ES 209	446873	2092.P20.gz43_274709
M00043490A:F03	ES 209	456577	2092.P24.gz43_274773
M00063157B:B09	ES 209	464791	2102.A16.gz43_275283
M00063165C:F10	ES 209	449206	2116.A13.gz43_306321
M00063457A:B12	ES 209	558147	2102.B18.gz43_275316
M00063493D:G07	ES 209	644927	2102.F08.gz43_275160
M00063496B:F07	ES 209	558900	2102.F14.gz43_275256
M00063507B;B08	ES 209	621081	2102.G10.gz43_275193
M00063514B:E04	ES 209	498509	2102.H02.gz43_275066
M00063546B:F01	ES 209	730600	2116.D21.gz43_306452
M00063552B:B06	ES 209	89082	2116.E15.gz43_306357
M00063580A:A07	ES 209	649744	2116.I06.gz43_306217
M00063592B:E09	ES 209	378453	2116.J09.gz43_306266
M00063601D:C05	ES 209	379154	2102.I24.gz43 275419
M00063803B:F11	ES 209	402353	2103.B13.gz43 275620
M00063861C:F09	ES 209	544229	2103.E13.gz43 275623
M00063864A:H10	ES 209	380514	2103.E17.gz43 275687
M00063865A:F01	ES 209	427540	2103.E21.gz43 275751
M00063866B:H02	ES 209	643609	2103.F01.gz43 275432
M00063900B:B04	ES 209	730845	2103.H23.gz43 275786
M00063923B:A04	ES 209	402941	2103.J21.gz43 275756
M00063927A:B08	ES 209	554395	2103.K10.gz43 275581
M00063943B:G12	ES 209	377696	2103.M06.gz43 275519
M00063943B:G12	ES 209	377696	RTA22200255F.f.15.1.P
M00063970A:D09	ES 209	642263	2116.M13.gz43 306333
M00063980B:A08	ES 209	378447	2116.N09.gz43_306270
M00063988A:C02	ES 209	404453	2116.O22.gz43 306479
M00063994B:D10	ES 209	535955	2116.P08.gz43 306256
M00064064D:D11	ES 209	554032	2104.A21.gz43 297640
M00064077B:H02	ES 209	639578	2104.C08.gz43_297434
M00064081A:D04	ES 209	52644	2104.C19.gz43 297610
M00064082C:D11	ES 209	505275	2104.D01.gz43 297323
M00064085B:A12	ES 209	649035	2104.D11.gz43 297483
M00064101B:E12	ES 209	376559	2104.E08.gz43 297436
M00064104B:A01	ES 209	558382	2104.E17.gz43 297580
M00064107C:E03	ES 209	463217	2104.E23.gz43 297676
M00064112A:G03	ES 209	646187	2104.F12.gz43 297501
M00064131B:A09	ES 209	647586	2104.H11.gz43_297487
M00064147B:G08	ES 209	454622	2104,I20,gz43 297632
M00064194B:A02	ES 209	11379	2104.N01.gz43 297333
M00064195C:B02	ES 209	446397	2104.N03.gz43 297365
M00064196D:C10	ES 209	469367	2104.N05.gz43_297397
M00064201A:C08	ES 209	727888	2104.N13.gz43_297525
M00064220B:E01	ES 209	418763	2104.P04.gz43_297383
L-100007220D.E01	202	410103	2104.1 04.8242_221303

Table 13

CloneID	ES No	ClusterID	SequenceName
M00064307C:E12	ES 209	404453	2117.D05.gz43_306580
M00064371B:E01	ES 209	728687	2117.J15.gz43 306746
M00064383B:B04	ES 209	551960	2117.K19.gz43 306811
M00064387A;H05	ES 209	447461	2117.L05,gz43 306588
M00064391C:D09	ES 209	471364	2117.L18.gz43_306796
M00064394A;C02	ES 209	416886	2117.M03.gz43_306557
M00064446D:C08	ES 209	446397	2118.A09.gz43_307025
M00064534D;H04	ES 209	730238	2118.H24.gz43_307272
M00064592D:F05	ES 209	177443	2118.M09.gz43_307037
M00064601C:H06	ES 209	644919	2118.N03.gz43 306942

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## We Claim:

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 An isolated polynucleotide comprising a nucleotide sequence which hybridizes under stringent conditions to a sequence selected from the group consisting of SEQ ID NOS: 1-6010.

- An isolated polymucleotide comprising at least 15 contiguous nucleotides of a nucleotide sequence having at least 90% sequence identity to a sequence selected from the group consisting of: SEQ ID NOS:1-6010, a degenerate variant of SEQ ID NOS:1-6010, an antisense of SEQ ID NOS:1-6010, and a complement of SEO ID NOS:1-6010.
- 3. An isolated polymucleotide comprising at least 15 contiguous nucleotides of a nucleotide sequence selected from the group consisting of: SEQ ID NOS:1-6010, a degenerate variant of SEQ ID NOS:1-6010, an antisense of SEQ ID NOS:1-6010, and a complement of SEQ ID NOS:1-6010.
- The isolated polynucleotide of claim 3, wherein the polynucleotide comprises at least 100 contiguous nucleotides of the nucleotide sequence.
  - The isolated polynucleotide of claim 3, wherein the polynucleotide comprises at least 200 contiguous nucleotides of the selected nucleotide sequence.
  - 6. An isolated polynucleotide comprising a nucleotide sequence of at least 90% sequence identity to a sequence selected from the group consisting of: SEQ ID NOS:1-6010, a degenerate variant of SEQ ID NOS:1-6010, an antisense of SEQ ID NOS:1-6010, and a complement of SEQ ID NOS:1-6010.
  - 7. The isolated polynucleotide of claim 6, wherein the polynucleotide comprises a nucleotide sequence of at least 95% sequence identity to the selected nucleotide sequence.
- The isolated polynucleotide of claim 6, wherein the polynucleotide comprises a
   nucleotide sequence that is identical to the selected nucleotide sequence.
  - A polynucleotide comprising a nucleotide sequence of an insert contained in a clone deposited as ATCC Accession No. PTA-2027, PTA-2028, PTA-2029, PTA-2030, PTA-2031, PTA-2032, PTA-2033, PTA-2034, PTA-2035, PTA-2036, PTA-2037, PTA-2038, PTA-2039, PTA-2040, PTA-2041, PTA-2042, PTA-2043, PTA-2044, PTA-2045, PTA-2046, PTA-2047, PTA-2050, PTA-2051, PTA-2052, PTA-2053, PTA-2054, PTA-2055, PTA-2056, PTA-2057, PTA-2058, PTA-

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2059, PTA-2060, PTA-2061, PTA-2062, PTA-2048, PTA-2049, PTA-2063, PTA-2064, PTA-2065, PTA-2066, PTA-2067, or PTA-2068.

- 10. An isolated cDNA obtained by the process of amplification using a polynucleotide comprising at least 15 contiguous nucleotides of a nucleotide sequence of a sequence selected from the group consisting of SEO ID NOS:1-6010.
  - 11. The isolated cDNA of claim 10, wherein the polynucleotide comprises at least 25 contiguous nucleotides of the selected nucleotide sequence.
  - The isolated cDNA of claim 10, wherein the polynucleotide comprises at least 100 contiguous nucleotides of the selected nucleotide sequence.
- The isolated cDNA of claims 10, 11, or 12, wherein amplification is by polymerase
   chain reaction (PCR) amplification.
  - 14. An isolated recombinant host cell containing the polynucleotide according to claims 1, 2, 3, 6, 9, or 10.
- 20 15. An isolated vector comprising the polynucleotide according to claims claims 1, 2, 3, 6, 9, or 10.
- A method for producing a polypeptide, the method comprising the steps of:
   culturing a recombinant host cell containing the polynucleotide according to claims claims
   1, 2, 3, 6, 9, or 10., said culturing being under conditions suitable for the expression of an encoded polypeptide;

recovering the polypeptide from the host cell culture.

- An isolated polypeptide encoded by the polynucleotide according to claims 1, 2,
   3, 6, 9, or 10.
  - 18. An antibody that specifically binds the polypeptide of claim 17.

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19. A method of detecting differentially expressed genes correlated with a cancerous state of a mammalian cell, the method comprising the step of:

detecting at least one differentially expressed gene product in a test sample derived from a cell suspected of being cancerous, where the gene product is encoded by a gene comprising an identifying sequence of at least one of SEQ ID NOS:1-6010:

wherein detection of the differentially expressed gene product is correlated with a cancerous state of the cell from which the test sample was derived.

- A library of polynucleotides, wherein at least one of the polynucleotides comprises the
   sequence information of the polynucleotide according to claims 1, 2, 3, 6, 9, or 10.
  - 21. The library of claim 20, wherein the library is provided on a nucleic acid array.
  - 22. The library of claim 20, wherein the library is provided in a computer-readable format.
  - 23. A method of inhibiting tumor growth by modulating expression of a gene product, the gene product being encoded by a gene identified by a sequence selected from the group consisting of SEO ID NOS:1-6010.